The topics covered with respect to chemical weathering on Mars include the following: Mars soil, mineralogy, spectroscopic analysis, clays, silicates, oxidation, iron oxides, water, chemical reactions, geochemistry, minerals, Mars atmosphere, atmospheric chemistry, salts, planetary evolution, volcanology, Mars volcanoes, regolith, surface reactions, Mars soil analogs, carbonates, meteorites, and reactivity. For individual titles, see N93-31934 through N93-31963.
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PREFACE

This volume contains papers that have been accepted for the Workshop on Chemical Weathering on Mars, September 10–12, 1992, in Cocoa Beach, Florida. The Program Committee consisted of Roger Burns (Massachusetts Institute of Technology) and Amos Banin (Hebrew University and NRC/NASA Ames Research Center).

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CONTENTS

The Mineralogy and Formation Processes of Mars Soil
A. Banin .......................................................... 1

Martian Weathering/Alteration Scenarios from Spectral Studies of Ferric and Ferrous Minerals
J. F. Bell III, J. B. Adams, and R. V. Morris ................................................................. 2

Effects of the Chemical Environment on the Spectroscopic Properties of Clays: Applications for Mars
J. L. Bishop and C. M. Pieters .......................................................... 4

Dehydroxylated Clay Silicates on Mars: Riddles about the Martian Regolith Solved with Ferrian Saponites
R. G. Burns .......................................................... 6

The Fate of Iron on Mars: Mechanism of Oxidation of Basaltic Minerals to Ferric-bearing Assemblages
R. G. Burns .......................................................... 8

Implications for Volcanogenic Volatile Release on the Weathering of Mars
B. C. Clark .......................................................... 9

Calculated Mineral Precipitation Upon Evaporation of a Model Martian Groundwater Near 0°C
J. D. DeBraal, M. H. Reed, and G. S. Plumlee ......................................................... 10

On the Weathering of Martian Igneous Rocks
G. Dreibus and H. Wänke .......................................................... 11

An Unusual Spectral Unit in West Candor Chasma: Evidence for Hydrothermal or Aqueous Alteration?
P. E. Geissler and R. B. Singer .......................................................... 12

Location of Nanophase Fe-Oxides in Palagonitic Soils: Implication for Martian Pigments
D. C. Golden, R. V. Morris, D. W. Ming, and H. V. Lauer Jr. ........................................ 14

Aqueous Geochemistry on Mars: Possible Clues from Salts and Clays in SNC Meteorites
J. L. Gooding .......................................................... 16

Effect of Purity on Adsorption Capacities of a Mars-like Clay Mineral at Different Pressures
T. Jenkins, B. McDoniel, R. Bustin, and J. H. Allton ........................................... 17
Low-Temperature Formation of Magnetic Iron-Oxides
C. B. Koch and M. B. Madsen ................................................................. 18

Implications of Isotopic Compositions of N and Ne in the Martian Atmosphere to Temporal Evolution of Its Atmosphere and Regolith
D. Lal ................................................................. 19

Preliminary Results of an Experimental Study of the Interactions of Basalt Glass and a Water Vapor Atmosphere: Implications for Weathering on Mars
J. J. Mazer, J. K. Bates, and C. R. Bradley .............................................. 19

Mars Brine Formation Experiment
J. M. Moore, M. A. Bullock, and C. R. Stoker ........................................ 20

Effect of DCB Extraction on Mössbauer and Spectral Data for a Hawaiian Palagonitic Soil (PN-9): Identification of Pigmentary Phases

Systematic Variations in the Spectral Properties of Bright Regions on Mars
S. Murchie, J. Mustard, J. Bishop, J. Head, C. Pieters, and S. Erard ................................................................. 23

Composition of Weakly Altered Martian Crust: Clues from Imaging Spectroscopy
J. F. Mustard, S. L. Murchie, S. Erard, and J. W. Head ................................................................. 25

Chemical Transport During Formation and Alteration of Martian Impact and Volcanic Deposits
H. E. Newsom ................................................................. 27

Reflectance Spectroscopy and GEX Simulation of Palagonite and Iron-rich Montmorillonite Clay Mixtures: Implications for the Surface Composition of Mars
J. B. Orenberg, J. Handy, and R. Quinn ................................................................. 29

Using the Viking Biology Experimental Results to Obtain Chemical Information about Martian Regolith
R. C. Plumb ................................................................. 30

Chemical Modeling Constraints on Martian Surface Mineralogies Formed in an Early, Warm, Wet Climate, and Speculations on the Occurrence of Phosphate Minerals in the Martian Regolith
G. S. Plumlee, W. I. Ridley, and J. D. DeBraal .................................................. 31
THE MINERALOGY AND FORMATION PROCESSES OF MARS SOIL. A. Banin, Hebrew University, Rehovot 76100, Israel and NRC/NASA Ames Research Center, Mail Stop 229-12, Moffett Field CA 94035, USA.

The mineralogical nature of Mars soil is far from being understood, nor are the formation time and weathering processes known. Quantitatively, the two major mineral-forming elements in Mars soil are silicon and iron, constituting 44% and 19% of the soil as SiO₂ and Fe₂O₃, respectively [1]. The silicate phases have been studied only briefly, mostly because of their limited spectral fingerprinting in the VIS and NIR. Much attention was given to the iron minerals in the soil, due to their pronounced absorption in the VIS and NIR, making them easily detectable by telescopic observations [2,3]. In the following, the available information on Mars soil mineralogy, mostly obtained by remote sensing, will be reviewed and it will be hypothesized that it leads to the suggestion that nanophase short-range-ordered (“amorphous”) phases of the silicates and iron oxides abound in the soil. This then raises several questions: Why are the major minerals in Mars soil poorly crystalline? What are the weathering pathways? Why did the secondary silicate and iron oxides in Mars soil not “ripen” and develop into well-crystallized phases despite the long period of time since aqueous weathering took place? Is it the kinetics? Antarctic weathering analogues do not support this. Alternatively, is it possible that the minerals in the thin top layer of Mars soil, which is what we have sampled and analyzed up to now, represent a relatively fresh and recent product of slow weathering processes of volcanic materials proceeding on the surface of Mars over hundreds of millions of years?

Earlier studies [4] have identified 60% “silicate” in the soil, using the Mariner 9 IRTM data. Montmorillonite (a smectitic aluminosilicate clay mineral) was suggested as a major component of the dust and soil on Mars, on the basis of more detailed analysis of the IRTM (Mariner 9) data [5]. Nonweathered basalt was rejected as it did not fit the spectral features. Nontronite and montmorillonite were initially suggested as the major silicate components on the basis of the elemental analyses of the soil [6]. This was supported by successful simulations of the Viking Labeled Release (LR) experiment using iron-enriched montmorillonite [7] and the similarity of this clay’s reflectance spectra in the visible range to those of Mars soil [8]. Another candidate was suggested, primarily on the basis of spectral telescopic data, to be palagonite, a weathering product of volcanic glass [9–12]. However, several palagonites tested did not simulate the reactivity of Mars soil as observed in the LR experiment [13]. A magnesium containing silicate (saponite?) was detected in the analyses of the NIR spectra at 2.2 μm [2]. However, the typical doublet of well-crystallized standard montmorillonite at 10μm was not observed in the IR spectra of Mars, leading to the conclusion that no crystalline smectite is present in the soil [3] or, alternatively, that it constitutes less than 15–20% of the soil [14]. If these observations are further corroborated, they suggest that most of the weathered silicates in Mars soil are of the low-crystallinity or amorphous varieties. No definitive identification of this phase was given until now. It is interesting to note in this context that a poorly crystalline silicate-aluminium mineraloid was detected in nonmetamorphosed inclusions in EET 79001, one of the SNC meteorites [15]. Furthermore, the possibility that the silicate “rust” (iddingsite) found in the interior of Nakhla is of preterrestrial origin has not been completely ruled out, since this SNC meteorite is a fall with very limited terrestrial weathering. The “rust” is a silicate alteration product produced perhaps by the action of fluids enriched in halides on olivine. The very limited data are consistent with di- or trioctahedral phyllosilicates with excess Fe and Si [16].

Iron, the second most abundant element in Mars soil, appears to be primarily present in its oxidized form (ferric), strongly suggesting chemical weathering of the basaltic parent material. The typical reflectance spectrum of Mars in the visible/short-wave NIR range [12,17,18] bears the strong fingerprinting of oxidized iron and has been reproduced more or less faithfully in laboratory measurements of many different iron-containing phases. They include “amorphous” iron oxides [19], palagonite [9,10], nanophase hematite deposited in silica matrix [11], and iron-enriched smectites [8,13]. One conclusion from the abundance of spectral analogues is that unambiguous identification of the iron mineral(s) in Mars soil, on the basis of reflectance only, is not possible at present. A more important conclusion is that the accumulating evidence shows that iron in the weathered component of Mars soil is mostly present in poorly crystallized clusters of oxyhydroxy ferric iron, or as crystalline iron oxides but in extremely small particle size ranges (“nanophases” or “nanocrystals”). Recent telescopic observations have detected typical hematite features at 860 nm, and possibly other crystallized iron oxides [12,20]. However, these features are very weak and estimates of the content of hematite and/or other crystalline phases are in the range of 1–3% or less. The content of the magnetic mineral in the soil is also estimated to be in the range of 1–3%. It may be maghemite, magnetite, or nanophase hematite that is typically superparamagnetic. However, the bulk of the iron oxide-oxyhydroxide in the soil is amorphous or short-range ordered and is characterized by extremely small particle sizes.

As a whole, the evidence now leads to the suggestion that much of the silicon- and iron-containing phases in Mars soil are short-range-ordered or “amorphous,” i.e., do not have well-developed crystallinity. This hypothesis is based on accumulating excellent telescopic observations during the late 1980s (1988–1990 Mars–Earth oppositions) and detailed laboratory studies on soil analogues. It opens a new and intriguing question—what is the peculiar mode of weathering that has produced these amorphous silicates and iron oxides? More specifically, what is the cause of the limited growth of crystals of secondary silicate and iron oxide minerals in the weathered soil on Mars. Given that currently the most accepted model for the soil formation is during earlier epochs (~3.5 b.y. ago) when Mars was “warm and wet” [e.g., 21,22], it is puzzling that only a minor portion of the silicates and iron oxides have crystallized and developed a more thermodynamically stable mineralogical composition and particle size distribution.

Similar silicate and iron oxide entities are found on Earth only as transitory phases that “ripen” rapidly (geologically speaking) to more stable minerals. Palagonites represent such a transitory assemblage of minerals produced as the initial weathering product of volcanic glass. The maximal age of palagonite deposits on Earth is in the range of 1-10 x 10⁶ yr.

Is it possible then that the peculiar nature of the martian dust and soil is due to a relatively “young” weathering product that has formed, and continues to form at an extremely slow rate, over the last several hundred million to a billion years? It is then plausible that this “recent” (in Mars timescale of changes) and nonevolved weathering product is coating or burying ancient, more-evolved weathered mineral assemblages that formed in the earlier “warm and wet” epochs of Mars. This suggestion apparently goes against the “common wisdom” on weathering. The low-temperature and arid conditions on Mars seem to preclude hydrolytic weathering as a significant process on its surface [22]. However, “surface weathering,” invol-
MARTIAN WEATHERING/ALTERATION SCENARIOS FROM SPECTRAL STUDIES OF FERRIC AND FERROUS MINERALS. James F. Bell III, John B. Adams, and Richard V. Morris, NASA/Ames Research Center, Moffett Field CA 94035, USA; 2Department of Geology, University of Washington, Mail Code AJ-20, Seattle WA 98195, USA; 3NASA Johnson Space Center, Code SN2, Houston TX 77058, USA.

In this abstract we review the major aspects of our current knowledge of martian ferric and ferrous mineralogy based on the available ground-based telescopic and spacecraft data. What we know and what we don’t know are used to constrain various weathering/alteration models and to identify key future measurements and techniques that can distinguish between these models.

What We Know: Ferric mineralogy. There is obvious and abundant evidence that oxidized Fe²⁺-bearing minerals exist on Mars. Spectroscopic observations and Viking data over the past 20 years have revealed that the ferric mineralogy occurs in two distinct styles: (1) poorly crystalline or even amorphous Fe³⁺-bearing materials that spectrally resemble certain terrestrial palagonites [e.g., 1-3]; and (2) well-crystalline ferric oxides like hematite (α-Fe₂O₃) or maghemite (γ-Fe₂O₃) [e.g., 4-6]. The available data indicate that the palagonite-like phases are spectrally dominant, and that the well-crystalline ferric oxides cannot constitute more than about 4-6 wt% abundance (if intimately mixed) [7]. All the noncondensable surface regions have a ferric spectral signature; however, the classical bright regions are distinctly “redder” than the classical dark regions and recent data have shown that the bright regions have generally enhanced crystalline ferric oxide absorption features [7]. Higher spatial resolution telescopic imaging and Viking Orbiter color data [e.g., 8] have revealed several small dark regions with increased crystalline ferric absorptions, though, and thus the albedo-crystallinity correlation is not globally uniform. Viking Lander color imaging reveals that the apparently unaltered rocks are covered to varying degrees by ferric-rich dust at the two locations studied [e.g., 9].

Ferrous mineralogy. There is much more debate and uncertainty about the occurrence and distribution of martian Fe˘-bearing minerals. Certain telescopic [1,10] and spacecraft [11,12] datasets have shown evidence for weak 1-μm and 2-μm Fe²⁺ absorption bands that preferentially occur in the dark regions, but other observations have either not clearly revealed such features [5] or have not seen a correlation between albedo and Fe²⁺ band depth [13]. In data where the band has been observed, it has been interpreted as evidence of low-Ca, high-Fe clinoxyroxene, indicating (in agreement with Viking Lander compositional results) that the martian “basement” rocks are mafic in composition.

Summary and other minerals. The only mineral definitively identified on the martian surface is hematite [4,5,7]. A possibly different poorly crystalline ferric-bear Initially there are two sets of low-resolution investigations per this study. The work reported here was supported in part by the NASA Exobiology Research Program and by grants from the National Research Council and the Hebrew University.

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Summary and other minerals. The only mineral definitively identified on the martian surface is hematite [4,5,7]. A possibly different poorly crystalline ferric-bear
phase itself. Pyroxene in some form probably exists [1,10], but conclusive mineralogic identification is still forthcoming [e.g., 12]. Data at longer near-IR wavelengths and simulations of Viking experiments have been used to claim the existence of iron-rich clay minerals [e.g., 15] and carbonate-bearing minerals like scapolite [16]. Confirmation of these mineralogic identifications by other researchers and other techniques has not yet been provided, though.

What We Don't Know: The presence of hematite masks the spectral identification of many other crystalline ferric oxide phases, like goethite (α-FeOOH), at telescopic spatial scales [7,17]. This provides a severe constraint on our current ability to identify almost all the ferric oxyhydroxides and iron-rich smectites in the currently available spectral data. Terrestrial analog studies have shown that several different extremely fine-grained ("nanophase") ferric oxides are spectrally consistent with the Mars data. These include nanophase hematite [3,4] and lepidocrocite [18]. In addition, the poorly crystalline mineral ferricyanide (a hematite precursor in many terrestrial weathering environments) is also spectrally consistent [7]. Thus, there is no current consensus on the mineralogy of the poorly crystalline ferric oxide phase.

The identification of the ferrous mineralogy is clouded somewhat by the presence of abundant and spectrally dominating ferric minerals. For example, there are several ferric oxyhydroxide minerals with weak absorption features in the 0.90-1.05-μm region [e.g., goethite, akaganeite (β-FeOOH)]. Hematite masks the shorter wavelength absorption features of such minerals, and thus it is not readily clear to what extent the surface ferric mineralogy may be enhancing the features from (or be mistaken for features due to) the surface ferrous minerals. In addition, the distinction in the data between well-crystalline ferrous minerals like pyroxenes and ferrous glasses and tephra (which also can exhibit a 1-μm absorption feature) has not been examined in detail.

Possible Weathering Scenarios Consistent with Available Data: Here, we define "weathering" as the steady-state oxidation and/or hydration of exposed surface materials, specifically leading to the conversion Fe2+ → Fe3+. This is to be distinguished from "alteration," which we define as a more forced process resulting from, for example, hydrothermal circulation, impact into ground ice, etc. The available data are consistent with two primary models for the weathering of martian surface materials.

Model 1. The exposed bedrock and surface rocks never have been weathered and are simply covered by and/or mechanically mixed with variable amounts of mobile Fe3+-bearing materials. This implies that these unweathered rocks are not the source of the global dust, and that the dust was formed in the past under different conditions. The currently exposed unweathered rocks could have escaped weathering by not being exposed during previous weathering events, or these rocks may postdate weathering conditions (e.g., later eruptions of basalt). The original source materials for the dust may currently be either buried or completely eroded. This model implies that there may be areas of oxidized (older) and unoxidized (postdust formation) bedrock materials on Mars.

Model 2. Exposed bedrock and surface rocks have been weathered in the past or are actively weathering, resulting in the creation of Fe3+-rich weathering rinds or cracks, and spallation or mechanical breakdown then frees the weathered materials. This implies that the dust is a direct product of the weathering of bedrock and/or tephra by surface reactions.

There are important climatic and mineralogic implications for each of these scenarios. If the exposed rocks are not actively weathering and producing the ubiquitous ferric-rich dust and soil (model 1), then the question must be asked, where did the weathering products come from? Two possibilities are immediately obvious:

(1a) Current reaction rates are exceedingly slow [21], leaving the exposed rocks in metastable thermodynamic disequilibrium, but reaction rates at some point(s) in the past were substantially higher, allowing oxidative weathering of bedrock/ephra to occur and to generate globally significant amounts of ferric material (i.e., model 2 was dominant in the past); or
(1b) A single event or series of events (e.g., impacts, intrusions, etc.) has produced all the observed weathering products over the course of martian geologic history [see 3].

Model 1a requires substantial climatic variations (increased T, P, RH) at some point(s) in the past. Model 1b could operate either in the current climatic regime or under warmer, wetter conditions. In either case, model 1 argues that the present inventory of surface weathering products is a fossil remnant from the past and thus the distribution of Fe3+-bearing phases may have little or no genetic relationship to the locally underlying bedrock.

If well-defined pyroxene absorption features occur in telescopic and/or spacecraft spectral data on 10-250-km spatial scales, then there must be exposures, at some subpixel scale, of relatively pure, unaltered pyroxene. Viking Lander images at Chryse and Utopia did not show any evidence that the ferric-rich soils and dust were genetically related to the abundant local rocks [9]. These arguments favor model 1. If, however, only weak, broad Fe2+ spectral features are observed, then it may be possible that the pyroxene (or other basaltic glass or minerals) are coated by a thin ferric weathering layer [19,20]. Telescopic and Viking observations of apparently ferric-rich dark regions (e.g., parts of Elysium Planitia [7,8]) provide some data that are more consistent with such an active weathering scheme (model 2).

What Needs To Be Done To Narrow The Possibilities: The key to understanding both the current and past martian weathering/alteration environment lies in being able to accurately measure the spectra of the rock "end member." If the spectral signature of the ferric-rich surface soil and dust (and of the intervening atmospheric dust) can be properly removed, then it will be possible to determine whether the underlying basalt is "lunar" (not at all oxidized) or "terrestrial" (actively oxidizing). This is an extremely difficult task in practice, because it requires simultaneous knowledge of the surface spectral reflectance (or albedo) and the airborne dust opacity, as well as various assumptions about (or direct measurements of) the dust composition and scattering properties. This is certainly hard to do at grounded telescopic spatial resolution, but is much more possible and practical using a Mars-orbiting platform.

Our understanding of the currently available spectral data can be substantially enhanced by additional experimental and theoretical work on mixtures of basalts and tephas, poorly crystalline ferric materials, and well-crystalline ferric oxides and oxyhydroxides. For example, as mentioned above, the identification of band centers and absorption band depths of Fe3+-bearing minerals in the 1-μm region may be influenced by the presence of certain crystalline ferric oxides/oxyhydroxides. Both intimate and areal mixture studies of the detectability of various ferrous minerals, including glasses, within a ferric background (and vice versa) are needed. We also need more experimental and theoretical data on the variability of spectral parameters such as band depth, width, and center position given different systematic mixtures of potential Mars minerals under varying degrees of atmospheric dust and condensate opacity. Finally, it may be possible to compare the results of other age-dating techniques (craters, superposition) with UV-visible spectral slope or "redness" derived from Mars Observer and/or ground-based telescopic measurements to determine whether or not the less-oxidized rocks are younger than the more oxidized rocks, dust, and soils.
EFFECTS OF THE CHEMICAL ENVIRONMENT ON THE SPECTROSCOPIC PROPERTIES OF CLAYS: APPLICATIONS FOR MARS. Janice L. Bishop and Carlé M. Pieters, Department of Geological Sciences, Brown University, Providence, RI 02912, USA.

Introduction: Laboratory studies of Mars soil analogs pose unique problems, since soils interact readily with their environment and exhibit variable characteristics depending on the environment. We have performed a series of experiments focusing on the spectral properties of clays and how they vary as a function of composition and environment, including examination of fundamental as well as overtone absorptions, that occur in the mid- and near-IR respectively. Smectitic clays have been selected in our laboratory experiments as a primary surface analog for Mars because of their compatibility with results of the Viking biology experiments [1], their stability under current martian conditions, and their compatibility with reflectance spectra of Mars [2]. We prepared a number of monoionic montmorillonites in order to examine the influence of cations on the water molecules in the clay interlayer region. Mössbauer spectra of several montmorillonites with variable amounts of interlayer iron confirm the presence of ferricyrite [3].

General Character of Clay Absorption Bands: Montmorillonite exhibits several strong features due to interlayer water, structural OH, and the silicate layers as shown by the composite RELAB spectrum in Fig. 1. Fundamental molecular water vibrations occur at ~3 μm (νw) and 6 μm (νOH), which have combination bands at 1.45 μm and 1.95 μm. Interlayer cation-bound molecular water has a stretching vibration (νw+) at a lower wavelength than that of the extensively H-bonded interlayer water (νOH), resulting in combination bands at ~1.41 μm and ~1.91 μm. Structural OH has fundamental vibrations at 2.75 μm (νOH) and ~11 μm (δM-OH, M = Al, Fe, Mg), a combination at 2.2 μm, and an overtone at 1.41 μm. Both the strength and energy of these absorptions are sensitive to the chemical environment of the OH and H₂O molecules. For instance, the polarizability of the octahedral and interlayer cations influences the OH and H₂O stretching and bending vibrational energies. One phase of our initial experiments has focused on the influence of interlayer cations on the character of near-infrared hydration bands [4]. Spectral characterization of clays and hydrated minerals is based on the H₂O and OH vibrational energies, which are dependent on the chemical environment of the bonds. The nature of interlayer water is affected by the interlayer cations and the degree of H-bonding.

Interlayer Cation-Water Interactions: Exchange experiments. In our laboratory experiments Fe²⁺, Mg²⁺, Ca²⁺, Na⁺, and H⁺ are exchanged for the naturally occurring mixture of Na and Ca interlayer cations in SWy-1 montmorillonite (obtained from the Clay Mineral Society, Source Clays Repository) using a procedure similar to that of Banin [2]. The amount of exchanged cation added was calibrated for 100% cation exchange capacity (CEC), which is the amount of interlayer cations required to compensate the charge deficit from octahedral layer substitutions. Fe²⁺-rich montmorillonites were prepared in a similar manner with up to 600% CEC.

Water analyses. The nature of interlayer water molecules in smectite clays is known to be quite complex [5]. Some H₂O is readily removable by mild heating and drying, while more resilient H₂O remains in the clay to above 300°C. Spectroscopic studies [4,6] suggest that water molecules can be present in many forms in the
interlayer regions: (1) strongly cation-bound, inner sphere H$_2$O, (2) strongly surface-bound H$_2$O, and (3) strongly H-bonded, liquid-like water. We have obtained chemical information about the form of water present in several homoionic clays through infrared spectroscopy and digital thermal analysis. We attribute the low temperature (<200°C) water loss to the liquid-like interlayer water, the water loss in the 350°C to 550°C range to inner sphere cation-bound water, and the water loss at 600°C to the structural OH. We have found for both dry and moist relative humidity environments that the amount of removable water in the montmorillonite samples varied depending on the interlayer cation (Na < Fe$^{3+}$ < Ca < Mg). This trend corresponds to adsorption isotherms for Na-, Ca-, and Mg-smectites [7].

Strength of bands. We found that the amounts of removable water in our suite of cation-exchanged clays correlates with the 1.9-μm and 6-μm band depths. The strengths of the 1.9-μm and 6-μm hydration features thus depend on the kind of interlayer cation, as well as on the relative humidity of the environment, whereas the strength of the 2.2-μm feature is primarily due to structural OH and octahedral cations. Reflectance spectra of natural (Na/Ca), Fe$^{3+}$, Ca-, and Mg-montmorillonite showing the relative strengths of the 1.9-μm and 2.2-μm hydration features are presented in Fig. 2.

The interpretation of Mars spectra serves as an example of the need for an improved understanding of the relationships between chemical environment, fundamental vibrations, and overtones of hydration features in clays. Some confusion over the presence of molecular water and structural OH in Mars surface material exists because a strong band has been observed at 2.7-3.0 μm (attributed to fundamental molecular water) and weak features have been suggested at 2.2-2.35 μm, but the characteristic water combinations and overtones at 1.4 μm and 1.9 μm are difficult to confirm. These features at 1.4 μm and 1.9 μm are composed of multiple overlapping peaks that vary as a function of octahedral cation, interlayer cation, and amount of interlayer water molecules [4]. An important application to Mars from our recent work is that a ferric montmorillonite in a dry environment would have a 1.9-μm absorption that is weaker than the 2.2-μm absorption, and thus the 1.9-μm absorption would be difficult to observe in Mars spectra.

Fe$^{3+}$-rich Clays and Ferricydrate Formation: Smectite clays readily incorporate aqueous ions and chemical complexes into their interlamellar regions. Clays are therefore highly sensitive to their chemical environment and clays on Mars should reflect the martian environment in the nature of their interlayer regions. In addition, smectite clays are able to serve as a template for chemical reactions in the interlayer regions [8], and interlayer expansion would physically allow production of nanophase products, but would restrict long-range order.

Several Fe$^{3+}$-rich montmorillonite samples were prepared containing 2-10 wt% (as Fe$_2$O$_3$) interlayer iron, in addition to the 4 wt% (as Fe$_2$O$_3$) structural iron already present in our Wyoming montmorillonite samples. During the exchange process Fe$^{3+}$ ions are added to the clay solution at pH 2, then the pH is increased to 7, causing the Fe(H$_2$O)$_6^{3+}$ complexes to polymerize to Fe$_3$(H$_2$O)$_6$(OH)$_2^{2+}$ and Fe$_2$(H$_2$O)$_6$(OH)(OH)$_2^{2+}$, losing H$_2$O$_2$s and H$_2$s as bridges are formed [9]. Concentrated Fe$^{3+}$ complexes react further, forming double chains of [Fe(O,OH,H$_2$O)$_4$] units and eventually ferricydrate [10,11]. Ferricydrate is thus predicted as an important step in weathering processes of ferric materials.

We have evaluated several of our montmorillonite samples using Mössbauer spectroscopy in a cryogenic facility at MIT in collaboration with R. Burns. Mössbauer studies of our Fe$^{3+}$-rich montmorillonites have produced magnetic hyperfine splitting patterns that confirm the presence of ferricydrate in these Fe$^{3+}$-rich clays [3]. Room-temperature spectra of all our montmorillonite showed a quadrupole doublet, characteristic of the structural iron in montmorillonite. At 4.2 K a sextet indicating magnetic ordering was observed for the Fe$^{3+}$-rich montmorillonites, but not the Na and Ca montmorillonites. This sextet was significantly stronger in the samples containing higher quantities of Fe$_2$O$_3$, and is characteristic of crystalline ferric oxides. The magnetic splitting in the Mössbauer spectra of our Fe$^{3+}$-rich clays is 48.5 T, which is within the range of 48.2-50.5 T for ferricydrate [12]. The magnetic splitting of other ferric oxides is measurably different, for example, hematite: 54.6 T; goethite: 50.6 T; and lepidocrocite: 45.8 T [10].

Furthermore, visible reflectance spectra of this Fe$^{3+}$-rich, ferricydrate-bearing montmorillonite are characterized by a weak, broad absorption at ~0.9 μm, a reflectance maxima at ~0.75 μm, an inflection point at ~0.6 μm, and a steep falloff toward ultraviolet wavelengths (Fig. 3). These features are not observed in natural (Na/ Ca), H-, Na-, Ca-, or Mg-exchanged SWY montmorillonites. As shown in Fig. 3, reflectance spectra of the Fe$^{3+}$-rich montmorillonites correspond remarkably well to telescopie data of several bright areas of Mars [3]. The Syria Planum spectrum was measured by Bell [13] and the Lunae Planum spectrum was measured by McCord [14].

Continuing Experiments: Although infrared reflectance spectroscopy is routinely used to study planetary surfaces and atm-

![Fig. 2. Reflectance spectra of homoionic montmorillonites (1.8-2.3 μm) (note relative strength of 1.9- and 2.2-μm absorption).](image-url)

![Fig. 3. Reflectance spectra of natural (Na/Ca) and Fe$^{3+}$-rich montmorillonites and telescopie spectra of Mars (0.3-1.2 μm).](image-url)
spheres, very little infrared research has been performed on the interactions between these surfaces and their atmospheres. We plan to use an environmental chamber to explore surface-atmosphere interactions that will provide information relevant to Mars. In order to better understand the nature of interlayer water in smectite clays and the cation-water interactions, we are planning spectral experiments for our homoionic (Na, Ca, Mg, Fe(3+)) montmorillonites that will (1) measure shifts in vibrational energy at 1.9 μm, 3 μm, and 6 μm at 25°C while varying the relative humidity of the sample chamber and (2) measure variations in the spectral character of these features as the temperature is elevated from 25°C to 300°C to study the clay hydration features in an ultradry environment. We are also designing experiments to measure spectra of the Fe(3+) rich montmorillonites at low temperatures (100–250 K), which will allow us to study the hydration bands in clay-ice systems.

In order to evaluate the behavior of clay systems under both current and more element possible past environments on Mars, we are currently planning aqueous and anhydrous reactions with sulfates, carbonates, and chlorides in Fe(3+)–rich montmorillonites. Spectra will be measured for these samples with a focus on the development of absorptions relating to Fe(3+)-sulfate, Fe(3+)–carbonate, and Fe(3+)-chloride bonds. We plan to measure spectra of these new and complex Fe(3+)–rich montmorillonites at cool temperatures (100–250 K) to study clay-ice minerals. Perhaps these experiments will lead to novel ideas for duricrust models. The nature of selected iron oxides and iron material formed in the interlayer region of these Fe(3+)-rich samples will be characterized using Mössbauer spectroscopy at room temperature and 4.2 K.


\[ \text{Fe}^{2+} + \text{OH}^- + 1/2 \text{O}_2 \rightarrow \text{Fe}^{3+} + \text{O}_2^- + 1/2 \text{H}_2\text{O} \]

resulting in the formation of a dioctahedral smectite intimately associated with the ferric oxide phase [12,15]. However, intracrystalline redox reactions involving the oxidation of Fe(2+) to Fe(3+) ions within the clay silicate structure is also possible, by analogy with processes leading to the formation of oxyphiboles and oxybiotites [16–18]. Two different oxidation mechanisms exist [19]. First, dehydroxylation by oxygenation may occur when the clay silicate is exposed to atmospheric oxygen or is heated in air (reaction (2))

\[ \text{Fe}^{2+} + \text{OH}^- + 1/2 \text{O}_2 \rightarrow \text{Fe}^{3+} + \text{O}_2^- + 1/2 \text{H}_2\text{O} \]
Second, dehydroxylation by dehydrogenation may occur when the clay silicate exists in a vacuum or is heated in an inert gas or vacuum (reaction (3))

\[
\text{Fe}^{2+} + \text{OH}^- \rightarrow \text{Fe}^{3+} + \text{O}_2^- + 1/2 \text{H}_2
\]

In both cases, the crystal structure remains intact (i.e., the mineral has not decomposed), but the OH groups have been replaced by O²⁻ ions.

Spectroscopic studies of phyllosilicates including saponites, based on changes of peaks in the 3600–3700 cm⁻¹ region [e.g., 18, 20–22], indicate that hydroxyl groups close to three Mg²⁺ in the two cis- and one transoctahedral sites are relatively stable. However, cation clusters such as MgMgFe²⁺ and MgFe²⁺Fe⁺ in these octahedral sites are vulnerable to dehydrogenation and oxidation resulting from oxidation of structural Fe²⁺ ions to Fe³⁺ (reactions (2) and (3)). Hydroxyl groups associated with vacancies, corresponding to cation clusters such as MgFe²⁺(V) and Fe²⁺Fe⁺(V), are particularly vulnerable to conversion to O²⁻ ions. Extensive intracrystalline oxidation of Fe²⁺ to Fe³⁺ produces additional spectral features attributable to hydroxyl groups adjacent to cation clusters such as MgFe²⁺Fe⁺, MgFe²⁺(V), etc. These clusters contribute weak peaks to poorly defined spectra in the fundamental (∼2.75 μm) and first overtone (∼1.41 μm) hydroxyl stretching regions. Similar effects may produce additional peaks in the 2.2–2.4-μm region in infrared spectra of partially dehydroxylated Fe²⁺Fe⁺ saponites, but these effects have not yet been studied in detail [23, 24]. However, they could be contributing shoulders to bands measured around 2.35 μm in remote-sensed reflectance spectra of the martian regolith[5, 25, 26].

**Terrestrial Occurrences of Ferromagnesian Smedrites:** On Earth, Fe²⁺ saponites have been identified in several environments, including weathered ultramafic rocks [27], gabbros [28], vitric tuffs [29], oceanic and continental basalts [30–32], and submarine depressions associated with brines and metalliferous sediments [15]. These ferroan saponites rapidly oxidize to mixed-smectite assemblages containing ferrous saponites, Fe³⁺-montmorillonite, and nontronite, which are often intimately intergrown with ferrhydrite, nanophase Fe₂O₃, or FeOOH. Some ferromagnesian smectites occurring in alteration products of oceanic and continental basalts appear to have compositions intermediate between di- and trioctahedral smectites. For example, a smectite from oceanic basalts [31], originally formulated as

\[(M^+)₄(Mg₀.76Fe²⁺₀.₂⁴Fe³⁺₀.₅₅)₂·₈₀(Si₃.₄₀Al₀.₅₉Fe³⁺₀.₀₁)O₁₀(OH)₂\]

has a total octahedral occupancy of only 2.80. Mössbauer spectral data suggesting that this smectite is partially dehydroxylated, were confirmed by chemical analysis that indicated a deficiency of OH⁻ ions. Recalculation of the formula yielded [12]

\[(M^+)₂(Mg₀.₈₀Fe²⁺₀.₄⁴Fe³⁺₀.₅₉Al₀.₁₄)₀₃₀(Si₃.₅₃Al₀.₄₆)O₁₀.₇₉(OH)₁.₂₂\]

in which the dehydroxylated smectite is perfectly trioctahedral with a total octahedral occupancy of 3.00. Similar dehydroxylated ferrian saponites might have formed on Mars.

**Dehydroxylated Mg–Fe Smectites on Mars:** The Viking XRF analyses [1], when expressed as modal proportions of ionic species [33], yield the following formula units (g.ion/kg soil)

\[
\begin{align*}
\text{SiO}_{₆} & = 1.75; \ \text{Al}^³⁺ = 1.35; \ \text{Fe}²⁺ = 2.18; \ \text{Mg}²⁺ = 1.75; \ \text{Ti}⁴⁺ = 0.07; \ \text{Na}⁺ = 0.65; \\
\text{Ca}²⁺ & = 1.00; \ \text{SO}_{₄}²⁻ = 1.20; \ \text{Cl}⁻ = 0.25
\end{align*}
\]

Although the Mg and Ca are traditionally associated with SO₄²⁻ in evaporite minerals on Mars [34], the majority of the Mg may be partitioned as trioctahedral (Mg–Fe saponite) coexisting with dioctahedral (Al–Fe montmorillonite), with each dehydroxylated smectite accommodating ferric iron.

Terrestrial environments producing ferromagnesian smectites, such as low-temperature alteration of basaltic rocks and hydrothermal reactions involving brine-saturated metalliferous sediments, also probably exist on Mars. Upon exposure to the martian atmosphere, the ferroan saponites would undergo aerial oxidation by reactions (1) and (2), with dehydroxylated smectites being formed by reaction (2). Restoration of OH⁻ by protonation of the O²⁻ is known to cause some of the Fe³⁺ ions to be expelled from the octahedral sites in the clay silicates [12], thereby creating vacancies in the phyllosilicate structure and the precipitation of ferrhydrite between the layers. Such a mechanism accounts for the suggested occurrence of ferrhydrite associated with clay silicates on Mars [35]. On the other hand, reversal of reaction (2) would liberate oxygen from the regolith, accounting for results obtained in the Viking gas exchange experiments [36].

Spectral features originating from MgMgMg, MgMgFe²⁺, MgFe²⁺(V), etc., cation clusters, as well as AlAl(V) and AlFe²⁺(V), are likely to contribute fine structure to peaks in the fundamental and first-overline hydroxyl stretching frequencies at 2.75 μm and 1.41 μm, as well as in the 2.2–2.4-μm region. Thus, features attributed to HSO₄⁻ or HCO₃⁻ in scapelite in remote-sensed reflectance spectra of Mars [26] may originate, instead, from partially dehydroxylated ferrian saponites in the regolith.

**References:**

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THE FATE OF IRON ON MARS: MECHANISM OF OXIDATION OF BASALTIC MINERALS TO FERRIC-BEARING ASSEMBLAGES. Roger G. Burns, Department of Earth, Atmospheric, and Planetary Sciences, Massachusetts Institute of Technology, Cambridge MA 02139, USA.

Introduction: Perhaps the most conspicuous indication that chemical weathering has occurred on the surface of Mars is the overall color of the Red Planet and the spectroscopic features that identify ferric-bearing assemblages in the martian regolith [1,2]. Apparently, Fe²⁺ ions in primary minerals in parent igneous rocks on the martian surface have been oxidized to ferric iron, which occurs in degradation products that now constitute the regolith. The mineralogy of the unweathered igneous rocks prior to weathering on the martian surface is reasonably well constrained, mainly as a result of petrographic studies of the SNC meteorites [3]. However, the alteration products resulting from oxidative weathering of these rocks are less well-constrained. Thus, a variety of candidate ferric-bearing materials have been suggested, ranging from poorly crystalline ferric oxides [4], such as nanophase hematite [5], to altered volcanic glass or palagonite [6,7], various clay silicates [8,9], and goessaniferous ferrhydrite-jarosite-clay silicate assemblages [10]. Moreover, the mechanism and rate of oxidative weathering of surface rocks on Mars are poorly understood [11], raising numerous questions of chemical kinetics about the nature and extent of chemical weathering processes on Mars.

For example, when did oxidative weathering reactions take place on Mars? Were reaction rates faster in the past and did they necessitate an aqueous phase? Is the oxidized regolith a fossilized remnant of past weathering processes, or are chemical weathering reactions still occurring in the frozen regolith? If so, are they seasonal? Are the ferric-bearing assemblages metastable and can they be characterized more rigorously? Some of these questions are addressed here by examining experimental data bearing on the mechanism of oxidation of ferromagnesian silicates to insoluble ferric-bearing phases, which include pathways of dissolution, ferrolysis, and hydrolysis reactions that may be appropriate to conditions on the surface of Mars.

Primary Rocks Subjected to Chemical Weathering: Mineral assemblages in SNC meteorites [3,12], particularly the shergottites (Shergotty, Zagami, EETA 79001, ALHA 77005), indicate that olivine (Fe₉₋₁₀Mg₁₀₋₉), low-Ca pyroxene (pigeonite, Fe₉₋₁₀Mg₀₋₉), and high-Ca pyroxene (augite, Fe₉₋₁₀Mg₀₋₉) are the predominant ferromagnesian silicates in basaltic rocks that were subjected to chemical weathering on Mars. Lower Fe²⁺ concentrations occur also in plagioclase feldspar and accessory sulfide and oxide minerals [12], although local bodies of massive iron sulfide deposits may be present near the surface of Mars [13]. These basaltic sulfide, olivine, and pyroxene minerals are particularly vulnerable to chemical weathering reactions, which in aqueous solutions proceed through dissolution, oxidation, and hydrolysis stages that are strongly pH dependent [14]. The eventual hydrolysis products of dissolved ferric iron also depend on the type of anions present in the solutions [15,16]. Thus, Fe(III) phases precipitated from SO₄²⁻ and Cl⁻-bearing solutions are most relevant to Mars if brines have existed there [17].

Dissolution Processes: Dissolution of ferromagnesian silicates is generally congruent under anoxic conditions, indicating that soluble Fe²⁺, Mg²⁺, Ca²⁺, and silica dissolve from olivines and pyroxenes at rates proportional to their concentrations in the silicate minerals [14]. Incongruent solubility of pyroxenes may occur initially, however, both as a result of preferential release of cations in M2 sites and armoring of crystal surfaces by protective layers of insoluble ferric oxides when dissolution occurs under oxidizing conditions. Solubility rates of Fe³⁺ ions decrease in the order olivine > pyroxene M2 > pyroxene M1 >> feldspars [11,14]. Dissolution rates of these minerals are highest in acidic solutions, decrease with rising pH, are least at near-neutral pH, and increase when the pH is in the alkaline range. Rates of dissolution are slow near freezing temperatures and in solutions of high ionic strength. Since low temperatures and, perhaps, brines may currently exist on Mars, dissolution rates of basaltic minerals are probably extremely slow on the present-day martian surface, but may have been much faster in the past when warmer climates prevailed, particularly if acid rain and acidic groundwater conditions prevailed [10,18].

Oxidation of Dissolved Fe²⁺: Ferrous iron released during chemical weathering of ferromagnesian silicate and sulfide minerals is metastable with respect to ferric iron when aerated groundwater exceeds pH 3.5 [10]. In near-neutral and alkaline solutions, dissolved Fe²⁺ ions formed from Fe²⁺ ions are removed almost immediately; the rates of precipitation of ferric oxide and oxyhydroxide phases are then controlled by the supply of Fe³⁺ ions by dissolution processes. Reaction rates for the oxidation of dissolved Fe²⁺ ions are strongly pH dependent [19,20] and are very slow in acidic solutions. There is a 10-fold decrease in oxidation rate for each temperature-interval decrease of 15°C. Rates also decrease with increasing ionic strength, and are slower in sulfate-bearing solutions than in chloride-bearing solutions. Therefore, in brine eutectics proposed for Mars [17] (which are themselves slightly acidic due to hydrolysis reactions) that have equilibrated with oxygen in the current martian atmosphere (at P₂O₅ = 10⁻⁵ bar), rates of oxidation of Fe²⁺ ions are estimated to be at least 10⁶ times slower than in laboratory experiments performed at room temperature on Earth [11].

Mechanism of Polymerization of Hydrous Ferrile Oxides: Once Fe³⁺ ions are produced in solution, they are highly vulnerable to hydrolysis to poorly crystalline ferric hydroxyl gels or ferrilydrite, which age to different crystalline phases, depending on the rate of oxidation of dissolved Fe³⁺ to Fe⁴⁺, the concentration of Fe³⁺ ions, the nature of anions present, pH, temperature, and time of aging [15,16]. For example, rapid oxidation of dissolved Fe³⁺ at near-neutral pH produces lepidocrocite (γ-FeOOH) [21], which may revert to maghemite (γ-Fe₂O₃) by dehydration [22]. Such maghemite is one of

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[Figures and tables not shown in this text representation.]
the candidate magnetic phases in the martian regolith [22,23]. Fast oxidation of Fe$^{2+}$ in strongly alkaline solutions, on the other hand, produces δ-FeOOH or ferroxyhyte (δ-FeOOH), the magnetic properties of which also make it a candidate magnetic phase that peralkaline phases exist on the martian surface [24].

Hydrolysis of FeCl$_3$ solutions initially yields akaganeite (β-FeOOH), which may subsequently transform to goethite or hematite [25]. However, ferrihydrite that forms in the presence of NO$_3^-$ or ClO$_4^-$ anions ages to goethite (α-FeOOH), hematite (α-Fe$_2$O$_3$), or a mixture of the two, depending on the pH and temperature. Hematite is favored at elevated temperatures, low water activity, and neutral pH, whereas goethite formation predominates at both low and high pH. The mechanism of precipitation and aging of ferrihydrite are controlled by the types of the complex ferric ions in solution [26], which are influenced by temperature and pH, as illustrated in Fig. 1. Thus, monatomic Fe$^{3+}$ ions predominate only in highly acidic solutions and some monomeric complex ions (e.g., [Fe(OH)$_2^+$] and [Fe(OH)$_4^-$] are also present. However, it is polymerization of dimeric complex ions (e.g., linear [Fe$_2$(OH)$_2$]$_2^+$) and cyclic (Fe$_2$(OH)$_2$)$_2^+$ clusters [27], which are the major species in solution above pH 2, that generates goethite, the crystal structure that consists of double chains of edge-shared [FeO$_4$]$_2^-$ octahedra. The [Fe(OH)$_4^-$] species prevails at higher pH and elevated temperatures (Fig. 1) and facilitates polymerization of face-shared octahedra, the key feature of the hematite structure, by deprotonation-oxidation reactions [28]. Such a mechanism leading to the production of nanophase hematite would occur on Mars when permafrost sublimed, brine euctectics evaporated, and clay silicates were desiccated on the arid martian surface.

Another important polymerization mechanism relevant to sulfate brine euctectics on Mars involves hydrolysis products of ferric iron in H$_2$SO$_4$ solutions, which are distinctly different from those of other inorganic acids [16,29]. At low pH, the [Fe$_2$(OH)$_2$] complex ion predominates (Fig. 1c) [26], so that in strongly acidic solutions hydronium jarosite, (H$_2$O)Fe$_3$(SO$_4$)$_2$(OH)$_6$, may be precipitated [31], particularly in the presence of Na$^+$ and K$^+$ ions. By analogy with terrestrial gossans, jarosite formed during sublimation or evaporation of sulfate-bearing groundwater could have persisted on the arid surface of Mars [10].

**Applications to Mars:** Chemical weathering on Mars appears to have occurred when acidic groundwater interacted with surface basalts. In anoxic conditions in the deep-weathering environment, dissolution rates of silicate and sulfide minerals were fast, yielding high concentrations of Fe$^{2+}$, Mg$^{2+}$, silica, etc., in the acidic groundwater [10,18]. Soluble Fe$^{2+}$ ions could persist indefinitely in cold, subsurface water and be transported around the planet, provided anoxic conditions prevailed. When exposed to the oxygenated martian atmosphere, slow oxidation of dissolved Fe$^{2+}$ ions to Fe$^{3+}$ ions would occur and the ferric iron, too, would remain in solution indefinitely provided the groundwater remained acidic. Eventually, hydrolysis of dissolved ferric ions to hydrous ferric oxide occurred. Incomplete aging of the ferrihydrite to Fe$_2$O$_3$, which is retarded by dissolved silica [31], provided a mechanism for producing nanophase hematite, which is believed to occur on Mars [5]. Carbonate mineralization was not possible on Mars while acidic groundwater prevailed. Such conditions have probably persisted by the condensation of volcanic gases, through oxidative weathering of sulfide minerals, and during oxidation of dissolved Fe$^{2+}$ ions.

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**IMPLICATIONS FOR VOLCANOGENIC VOLATILE RELEASE ON THE WEATHERING OF MARS.** Benton C. Clark, Planetary Sciences Lab (B0560), Martin Marietta, Denver CO 80201, USA.

Volcanism on Mars has been widespread in both space and time [1]. Notwithstanding important specific differences between the mantles of Earth and Mars, the similarities are such that the suite of gases emitted from martian volcanic activity surely includes gases such as H$_2$, CO, CO$_2$, S-containing gases (H$_2$S, SO$_2$, or SO$_3$), and Cl-containing gases (e.g., Cl$_2$ or HCl). Both H$_2$O and CO$_2$ are present in the atmosphere of Mars; both are also present as surface condensates. Spectroscopic observations of the martian atmosphere clearly show that the S- and Cl-containing gases are severely depleted [2,3]. Likewise, there is no evidence of surface condensates of compounds of these elements. Within the soil, there is direct evidence of incorporation of H$_2$O [4] and some compounds of sulfur and chlorine [5]. None of the resultant weathering products have been directly identified, but both clays and salts have been indirectly implicated.

Estimates for the total magma generated after initial crustal formation and the terminal stages of heavy bombardment have recently been updated. Although some 60 times lower than the current areal rate of extrusive lava resurfacing on Earth [1], the martian total is nonetheless equivalent to the release of $\geq 500$ g/cm$^2$ of volatile averaged over the planet for every 0.1% wt/wt of volatile species released from magma. Quantitatively, this is sufficient to comprise as much as 5% of fine-grained regolith weathering product to a mean depth of 100 m. A reservoir of extremely fine-grained weathering product can be readily mobilized globally by the episodic dust storm activity. The relative absence of soil-consolidation factors, such as reworking by liquid water, tectonically driven metamorphism, and ubiquitous burial processes on Earth may permit survival of some material over significant portions of geologic time on Mars. As fines are repeatedly transported over the surface, they sample weathering regimes from various geologic settings and epochs.
These fines therefore could represent an undecipherable melange of past weathering processes, but also a planetwide sampling of the physical and chemical products from various surface materials. Excess acidity in the fines can occur due to the preponderance of acidic volcanic emissions. Some minerals will be more susceptible to attack than others [6]. The degree to which initial weathering reactions form barriers to further conversion of source material will be functionally dependent upon martian ambient conditions. Considerable interest would then center on the potential for comparisons between secondary products found as weathering rinds, vugs, or veins in their source rock and these eolian-transportable weathering product, presumably older and more reworked.

Resistant units such as large rocks and bedrock outcrops would be subjected to a balance between surface chemical weathering and physical removal by eolian abrasion. Because of saltation heights and wind shadowing effects, important three-dimensional geochemical gradients of secondary alteration products could be present in weathering rinds on large boulders such as those observed at the Viking 1 lander site.

Although carbonates and nitrates are widely expected in the martian regolith by many planetary geologists, current evidence is lacking or weak. Reworked fines may be scrubbed of any weathering product of either class of compounds since it has been demonstrated experimentally that volcanogenic SO₂ gas can undergo rapid heterogeneous phase displacement reactions with susceptible solid substrates even under simulated dry, cold martian conditions, releasing CO₂ and NO₃ respectively. The problem requires detailed modeling of functional relationships between quantities for each species released, reaction rates, and availability of labile substrates.

On the other hand, magmas release additional volatiles, many of higher atomic weight. Although it is experimentally difficult to identify and quantify all such emissions, a variety of data relevant to volatility would imply that weathering products may be highly enriched in elements such as Na, Cu, Zn, As, Se, Br, Rb, Cd, In, Sn, Sb, Hg, Y, Pb, and Bi compared to rock compositions [7]. Many of the compounds formed by these elements may be soluble in H₂O, as data indicate for the S- and Cl-bearing compounds in martian fines, and hence be subject to transport processes that create duricrust and soil pods.

Where weathering products have been exposed to bulk liquid water, chemical sediment deposits with complex evaporite sequences should be found on Mars. Quasitable liquid brine pools might also have resulted. However, if the dominant soluble anion on Mars is SO₄²⁻, then most strong freezing-point depressants salts would not be available for contemporaneous brine.

Although it is widely believed that the missing H₂O is buried in the regolith as physical deposits of permafrost ice, it cannot be ruled out that significant quantities, perhaps most, of this inventory have been incorporated into secondary minerals. The extent to which a pervasive drawdown of atmospheric volatiles has been mediated by chemical reaction with surface materials is important to understanding the past climatological history of Mars.


CALCULATED MINERAL PRECIPITATION UPON EVAPORATION OF A MODEL MARTIAN GROUNDWATER NEAR 0°C. J. D. Debraal, M. H. Reed, and G. S. Plumlee, Department of Geological Sciences, University of Oregon, Eugene OR 97403, USA. U.S. Geological Survey, Box 25046, Mail Stop 793, Denver Federal Center, Denver CO 80225, USA.

Previously we calculated [1] the effect of weathering a basalt of Shergotty meteorite composition [2] with pure water buffered at martian atmospheric values of CO₂ and O₂, to place constraints upon the composition of martian groundwater, and to determine possible equilibrium mineral assemblages. Figure 1 shows a revised calculation of the composition of the aqueous phase in the weathering reaction as a function of the amount of basalt titrated into the solution. The concentrations of sulfate and chloride ions increase in the solution from high water/rock ratios (w/r) on the left to low water/rock ratios on the right, until at w/r = 1, where 1 kg of basalt has been titrated, sulfate concentration is 1564 ppm and chloride is 104 ppm. This resulting fluid is dominated by sulfate and sodium, with bicarbonate and chloride at about the same concentration. We have evaporated this solution in an attempt to determine if the resulting evaporate can explain the Viking XRF data.

We used the program CHILLER [3] to evaporate this solution at 0.1°C. The solution was buffered at martian atmospheric values of CO₂ and O₂ (CO₂ = 10⁻²² bar, O₂ = 10⁻⁶ bar). Prior to evaporating the fluid, it was fractionated from its host rock to prevent back-reaction, under the assumption that the kinetics of evaporite precipitation are faster than those for solution reequilibration with its host rock. Evaporation was simulated by removing pure water from the

Fig. 1. pH and total modalities of aqueous components plotted vs. log grams of basalt added to pure water buffered by the martian atmosphere. Notice the accumulation of aqueous sulfate by oxidation of sulfide from the rock.

Fig. 2. Mineral abundance diagram that shows the presence and absolute abundance (in moles) of the evaporite minerals produced as 1 kg of martian groundwater is evaporated (see Fig. 3 for scale). Mineral abbreviations: dol - dolomite; qz - quartz; gyp - gypsum; daw - dawsonite; chl - chlorite; hem - hematite; non - nontronite.
Fig. 3. The pH and total modalities of aqueous components vs. percent martian groundwater evaporated.

Fig. 4. Detailed mineral abundance diagram for the last 1% of evaporation. The transition from gypsum to anhydrite occurs just after 98.7% evaporation. Mineral abbreviations: gyp - gypsum; anh - anhydrite; dol - dolomite; qz - quartz; chl - chlorite; micr - microcline.

system until activity coefficients could no longer be reliably calculated (after 99.8% evaporation). These results are plotted in Figs. 2 and 3.

Dolomite, quartz, dawsonite (NaAlCO₃OH₂), and nontronite saturate at the beginning of the evaporation and little change occurs in either the solution or the initial mineral assemblage until just after 65% of the water has been evaporated. At this point gypsum precipitates from the solution and becomes the dominant mineral in the assemblage. Gypsum is replaced by anhydrite once 99.8% of the water has been evaporated (Fig. 4), at which point the mineral assemblage is 84.7 wt% anhydrite, 15.1 wt% dolomite, with small amounts of quartz, chlorite, and hematite making up the difference. This assemblage is sulfur-rich (19.9 wt%), and if appropriately mixed with rock fragments from other sources, is consistent with the 3-4 wt% sulfur detected by the Viking XRF [4].

No chloride minerals precipitate in the evaporation of this fluid. At the point of anhydrite saturation, saturation indices (log Q/K) for halite and sylvite are ~1.65 and ~2.20, indicating that they were still significantly undersaturated. The Viking XRF data indicate that up to 0.7 wt% chloride is present in martian fines. Our results suggest that chloride may not reside in alkali halides, unless soil waters evaporate to dryness. An alternative residence for chloride may be in phosphates [5].


ON THE WEATHERING OF MARTIAN IGNEOUS ROCKS.
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Besides the young crystallization age, one of the first arguments for the martian origin of SNC meteorites came from the chemical similarity of the meteorite Shergotty and the martian soil as measured by Viking XRF analyses [1]. In the meantime, the discovery of trapped rare gas and nitrogen components with element and isotope ratios closely matching the highly characteristic ratios of the Mars atmosphere in the shock glasses of shergottite EETA 79001 [2,3] was further striking evidence that the SNCs are martian surface rocks.

The martian soil composition as derived from the Viking mission, with its extremely high S and Cl concentrations, was interpreted as weathering products of mafic igneous rocks. The low SiO₂ content and the low abundance of K and other trace elements in the martian soils point to a mafic crust with a considerably smaller degree of fractionation compared to the terrestrial crust. However, the chemical evolution of the martian regolith and soil in respect to surface reaction with the planetary atmosphere or hydrosphere is poorly understood. A critical point in this respect is that the geochemical evidence as derived from the SNC meteorites suggests that Mars is a very dry planet that should have lost almost all its initially large water inventory during its accretion.

In Shergotty, a H₂O content of 180 ppm was measured, compared to the about 2000 ppm H₂O in ocean ridge basalts. These 180 ppm H₂O correspond to 36 ppm H₂O for the martian mantle [4]. A mantle concentration of 36 ppm water would correspond to an ocean covering the whole planet to a depth of 130 m. According to all estimates, the degassing efficiency on Mars is thought to be small and outgassing of the interior is likely to have provided no more than about 10 m of water to the surface. These estimates are in conflict with the geological evidence for large amounts of water at the martian surface as derived from the large flood features and the valley networks [5].

The contradictory evidence of a dry martian mantle and the erosional martian surface features has been discussed by Carr and Wänke [6,7]. Recently, it has been found that part of the water present in SNC meteorites differs drastically in their oxygen isotopes from the oxygen bound in silicates [8]. This indicates that the martian surface water of today cannot be derived from the martian mantle in a direct way.

We do not have a definitive answer about the amount of water at the surface, but we have good reasons to assume that this water was very salty. The halogen content of the martian mantle as estimated from the chemistry of shergottites must be a factor of 3 higher compared to the Earth [4]. On today’s Earth most of the total water content is in the oceans, which have an average NaCl concentration of 3.1%. Hence, we should expect that the small amount of water on Mars was saturated with NaCl [9] at least at the time close to the final disappearance of liquid water on the martian surface. In addition to NaCl, MgSO₄ and MgCO₃ [10] are also expected to be present in the martian water. Recently, Wänke et al. [11] emphasized the importance of SO₃ on Mars. If we take 180 ppm H₂O in Shergotty at face value and compare it with the 1330 ppm S and 620 ppm C in Shergotty, it is evident that sulphur and carbon, respectively SO₂ and CO₂, are the most abundant volatiles in the Shergottya magma. Under martian surface conditions, SO₂ will be transformed quickly to SO₃, which in turn will react with H₂O to form sulfuric acid. Hence, it seems likely that in addition surface water became acidified.
To study the interaction between a fluid phase and martian surface samples we carried out leaching experiments with water and acids on the basaltic meteorite Shergott, the komatitite shergottite ALHA 77005, and the augite-olivine cumulate Nakhl. Leaching experiments on shergottites with brine water are in progress.

Nakhl contains alteration products of probably extraterrestrial origin that may be produced by the action of fluids rich in halides [12]. This "rust" contains high amount of Cl and K. The leaching experiment on Nakhl showed that of the very high concentration of Cl (1145 ppm) and Br (4.1 ppm) 96% resp. 50% are water-leachable, together with about 16% of the bulk Na and K, corresponding exactly to the leachable Cl as NaCl and KCl. Also sulphur is in Nakhl to 77% water-leachable. The water-leach was followed by an acid-leach (1N HNO₃). The treatment with this acid dissolved 80% of La, but only 16% Dy, 50% U, and 26% K. The La/Dy ratio of the leach is with 9.2 a factor of 3 higher than the La/Dy ratio in the bulk sample, but matches exactly the La/Dy ratio of the low-density fraction separated by Nakamura et al. [13]. However, these authors could not precisely identify this low-density fraction.

The leach experiments on the two shergottites show other features. In Shergott only 19% of the Cl and 15% of the Br was found to be water-leachable. At the following acid leach (1N HCl) the residual halogens were completely dissolved together with the whole phosphate-phase, whereas the treatment of ALHA 77005 with 1N HNO₃ leached halogens and phosphates from the bulk sample to only 70% [14]. This could mean that the sample was not sufficiently fine-ground or that part of P and the halogens are in insoluble phases, most probably glass.

We observed that in all acid-leach experiments of Nakhl and the two shergottites REE as well as U are considerably more leachable than K. The K/U ratio in Nakhl leach was 5400 compared to the residue ratio of 42,000. In Shergott leach a K/U ratio of 780 was found compared to a ratio of 37,000 in the residue. The reason for the difference of the behavior of K and U during weathering of martian rocks clearly lies in the fact that U as well as many other LIL elements, but not K, reside in the phosphates (apatite and whitlockite), which dissolve easily in dilute acids. The observation that acid weathering of igneous martian rocks yields drastically different K/U between leach and residue phases might be highly significant in respect to the interpretation of the K/U ratios derived from γ ray experiments by Phobos 2 and Mars 3 [15].


AN UNUSUAL SPECTRAL UNIT IN WEST CANDOR CHASMA: EVIDENCE FOR HYDROTHERMAL OR AQUEOUS ALTERATION? P. E. Geissler and R. B. Singer, Planetary Image Research Laboratory, Lunar and Planetary Laboratory, University of Arizona, Tucson AZ 85721, USA.

A spectrally distinctive unit on the floor of W. Candor Chasma (6°S, 76°W) in the central Valles Marineris may be a likely candidate for hydrothermal or aqueous alteration. This unusual material is noticeably redder than nearby plains and canyon floor-covering deposits of similar brightness in several Viking Orbiter color-composite images calibrated using PICS Level 1 procedures [1]. The surrounding plains and canyon floor units have colors that are typical for much of the weathered soil on Mars. Relative to adjacent materials, the West Candor unit has lower green-filter reflectance and higher red-filter reflectance (Fig. 1). While subtle, these spectral characteristics have been observed for this unit in a number of multispectral images acquired at different seasons and phase angles. When the color image data are transformed to hue, saturation, and value coordinates, the West Candor material stands out prominently among the rocks and soils of Coprates Quadrangle as a spatially coherent unit with a unique hue (Fig. 2). Physically, this means that the unit is compositionally distinct (unlike most of the bright materials in the region), since its spectral reflectance cannot be obtained through simple multiplicative scaling of the reflectance of the surrounding bright materials or by altering the spectral reflectance of the surrounding materials with an additive constant that is independent of wavelength. Recent studies of the directional reflectance properties of the surface in this region [2] indicate that the photometric phase function of the unit is similar to that of surrounding bright materials (isotropic to slightly backscattering).

In Viking high-resolution images (e.g., Fig. 3) the anomalous unit can be seen to correspond to two topographic depressions in heavily eroded floor-covering deposits in West Candor. Its restricted spatial distribution suggests that the unit may be genetically associated with the depressions. It is not clear whether we are seeing exposed underlying deposits distinct in composition, or later material deposited in preexisting depressions. Within these depressions are fine horizontal laminations possibly due to layering. However, Komatsu et al. [3] make a case that many of the apparent layers in the Valles

![Fig. 1. Comparison of West Candor Unit 3-point spectrum (solid line; 25 pixel avg.) with spectrum of nearby canyon wall rock of similar brightness (dashed line; 36 pixel avg.). Data from calibrated image composite, Viking Orbiter 1, orbit 383.](image-url)
Marineris interior deposits, including these, resemble erosional terraces and wave-cut platforms of terrestrial lakes and playas. Such erosion could have produced the morphology seen in West Candor even in unstratified, homogenous materials. In any case, this unit is apparently compositionally unique. The absence of similar material in depressions elsewhere in the canyon system seems to require a local mechanism for formation and/or alteration.

The Viking Orbiter bandpasses do not provide enough spectral information to generally allow unique mineralogic interpretations. Nevertheless, we think that the observed spectral reflectance characteristics of the West Candor unit are most readily explained by a composition somewhat richer in bulk crystalline hematite than typical for weathered or altered soils on Mars [e.g., 4,5]. Crystalline hematite, which is visually red, displays a strong and diagnostic crystal-field absorption near 0.53 µm [e.g., 4,6] to which the Viking green bandpass is quite sensitive despite that filter’s width. While crystalline hematite is not the primary coloring agent for typical soils on Mars there have been a number of observations consistent with its occurrence on the planet [4,7–10]. We interpret the Candor units to have a greater hematite content than typical martian soils, but to also contain the ubiquitous poorly crystalline or nanophase ferric oxide phase that gives Mars its characteristic orange-brown color.

Hematite is the most thermodynamically stable ferric oxide mineral under current martian surface conditions [11,12]. Crystalline hematite on Earth forms in a variety of environments through a number of mechanisms, including subaerial weathering of soils, hydrothermal alteration associated with volcanism or circulating mineralized fluids, and conversion of other previously formed hydrated ferric oxide minerals such as ferrihydrite and goethite. All these mechanisms might be viable on Mars as well, making it difficult to pinpoint the genesis of hematite in this Candor unit. The most common characteristic of hematite formation, relative to other ferric oxides, is heat. Increased temperature favors hematite, as does decreased water content and relatively neutral pH [e.g., 11,13]. The Candor units may have formed by primary volcanic oxidation, or through the action of circulating hydrothermal fluids. They might also have formed through aqueous precipitation of ferrihydrite in receding lakes, consistent with the localized nature of these deposits and the possibility of wave-cut terraces. This ferrihydrite could have subsequently converted to hematite through drying and application of heat. Whatever the origin, this unit is apparently unique, and required a geologically localized source of heat greater than current surface conditions, and greater than that to which surrounding weathered soils have been exposed. Localized volcanism, in one form or another, is a plausible source of that heat (cf. [14]).

We recommend that this region in Candor Chasma be observed carefully with upcoming missions to Mars. From Mars Observer, high-resolution images with the MOC should help resolve details of geologic relationships, including the question of layering vs. terracing. Data from the TES infrared spectrometer will be useful to search for possible clay minerals and/or evaporite salts. Unfortunately,
there is no visible-region spectral capability on Mars Observer to help refine our understanding of the ferric-oxide mineralogy. Mars 94 will carry two such instruments, the Omega visible and near-IR imaging spectrometer, and the Svet visible and near-IR imaging spectrophotometer. While we have not found other similar color units in the Valles Marineris region, they may exist elsewhere on Mars. Searches for such units throughout the large Viking Orbiter multispectral dataset will continue to be important.


LOCATION OF NANO-THE FE-OXIDES IN PALAGONITIC SOILS: IMPLICATION FOR MARTIAN PIGMENTS. D. C. Golden1, R. V. Morris1, D. W. Ming1, and H. V. Lauer Jr.2, INASA Johnson Space Center, Houston TX 77058, USA, 2Lockheed ESC2, Houston TX 77058, USA.

Introduction: Palagonitic materials from Mauna Kea, Hawaii, have been identified as Mars analogues based on their spectral and magnetic properties. These materials probably resulted from hydrothermal alteration during eruption of the volcano and/or from weathering under ambient conditions. The reflectance spectra of the Mars surface obtained by Earth-based telescopes and the reflectance spectra of analogues obtained in the laboratory show features due to electronic transitions of Fe(III) in oxide particles that range in size from nanometer (nanophase) to micrometer sized or larger. The presence of Fe(III) suggests oxidizing conditions during the alteration process in Mars that may have occurred in the past or during a slow ongoing process. In this study we examined two naturally altered basaltic samples from Hawaii (HWMK12 and HWMK13) and a laboratory-altered (PH-13-DCG2) basaltic glass similar in elemental composition to the above two samples. All three samples exhibited spectral characteristics similar to martian bright-region spectra. Chemical and mineralogical changes occurring at the surface of these basalts were studied in order to understand the basis for their Mars-like properties. The spectral properties of the three samples were examined after the removal of Fe oxides by chemical extractants.

Materials and Methods: Samples. Palagonitic soils HWMK12 and HWMK13 were collected from near the summit of Mauna Kea, Hawaii [1]. The <20-μm fraction of the HWMK12 and HWMK13 samples was used for all the investigations reported here. Sample PH-13, which is a black, relatively unaltered material, was obtained from the Puu Hulululu cinder cone ~15 km due south of the Mauna Kea summit by J. Bell. A 1-g sample of the <1-mm fraction of PH13 sample was heated in a teflon-lined vessel in the presence of 7 ml of H2O2, 4 ml of 1 M NaOH, and 5 ml of water for seven days at 200°C. The product (PH13-DCG-T2) was washed four times with deionized water, freeze dried, and stored in an air-tight bottle for further use.

Chemical extractions [2]. Acidified ammonium oxalate (pH = 3) (hereafter referred to as oxalate) was used to extract poorly crystalline Fe-oxides. A 0.3-M solution of ammonium oxalate was adjusted to pH 3 using 0.3-M oxalic acid solution. Extraction was performed by shaking 0.5 g of soil with 200 ml of the reagent in a polypropylene tube in the dark. A citrate-dihionate-bicarbonate (DCB) method was used to extract free Fe(III)-oxide. A solution of 0.3 M sodium citrate was mixed with 1 M NaHCO3 solution in the ratio 8:1. The pH was adjusted to 7.3 by adding 0.3 M citric acid. Extraction was performed by adding 25 ml of the reagent to 200 mg of the sample in a polypropylene tube at 75°C in a water bath followed by 1-g additions of sodium dithionite powder at 5-min intervals as necessary.

Untreated and extracted samples were analyzed by Mössbauer spectroscopy, reflectance spectroscopy, electron microscopy (SEM and TEM), and electron probe microanalysis.

Results and Discussion: The spectral data for natural samples HWMK12 and HWMK13 (<20 μm) are both characterized by relatively featureless absorption edges (Fig. 1). Compared to martian bright region spectra, the edges occur at somewhat shorter wavelengths. The bands near 1.4 and 1.9 μm are due to absorbed water on phyllosilicate surfaces or in zeolite structures. Both samples behaved in a similar fashion with respect to their reflectance spectra upon oxalate and DCB treatment. Upon oxalate treatment there is a slight shift of the absorption edge toward the shorter wavelengths and also a slight decrease in reflectance at longer wavelengths. The DCB treatment, however, had an appreciable effect (Fig. 1). The strength of the ferric absorption is reduced and there is a relative reflectivity maximum near 0.8 μm. The reflectivity maximum is the region between ferrous bands near 1.0 μm and ferrous-ferric charge transfer bands near 0.65 μm. The bands near 1 μm are also evident in the untreated samples.

The Mössbauer spectra of all three untreated samples are dominated by a ferric doublet that we have attributed to nanophase oxide (np-Ox) particles. The absolute intensity of the doublet did not decrease as much as expected on the amount of Fe extracted (Table 1). Apparently Fe and other elements were removed by the extraction procedures in a way such that the Fe content remained nearly constant.

Transmission electron microscopy of palagonitic soil sample HWMK12 before chemical treatment (Fig. 2a) shows the weathering basaltic glass matrix. The hydrated glass shows signs of polymerization of [Si(OH)4] increasing the porosity of the glass. Fe(II) in the original glass structure is oxidized and hydrolyzed as channels are formed during hydration and repolymerization of silica. Once a nucleus of Fe(OH)3 is formed Fe(II) diffuses in that direction and the iron oxide particle grows perhaps by surface catalyzed oxidation of Fe(II) near the original nucleus. As weathering proceeds, soluble cations and Si move away from the weathering front.

| TABLE 1. Fe-removal by oxalate and DCB extractions as a percent of the total Fe. |
|-----------------|-----------------|
| Sample          | Oxalate-Fe %    | DCB-Fe % |
| HWMK12          | 12              | 16       |
| HWMK13          | 9               | 17       |
| PH-13-DCG-T2    | ND              | 17       |


Fig. 1. The reflectance spectra of Mars analogue samples: (a) HWMK12, (b) HWMK13, and (c) PH-13-DCG-T2. Numbers 1 through 3 represent curves corresponding to untreated, oxalate-treated, and oxalate-DCB-treated samples respectively.

Fig. 2. Transmission electron micrographs of micromilled thin sections of HWMK12 sample mounted on holey carbon substrate showing two different weathering stages of basaltic glass: (a) initial glass repolymerization and Fe oxide nucleation, and (b) the accumulation of nanophase Fe oxide particles near the glass weathering front.

while leaving a residue of Fe oxides (the np-Ox particles). These cations and the silica precipitate as smectite or zeolite near the weathering front. Fe concentrations during hydration and oxidation of the glass can be seen as electron dense (dark) spots in the weathered zone (Fig. 2b). As the weathering advances, more ordered Fe oxide minerals are formed. Most of these oxides are still surrounded by the silica derived from weathered glass, which protects the Fe oxides from reducing agents like DCB. Therefore most of the np-Ox particles are not dissolved by oxalate or DCB treatment. Only the np-Ox particles on the surfaces are removed. The mineralogy of the hydrothermally altered PH-13 sample consisted of smectite, zeolite, and Fe-oxides as inferred from TEM, SEM, and X-ray diffraction.

Hydrothermal alteration in the laboratory of basaltic glass (PH-13-DCG-T2), similar in elemental composition to HWMK12 and HWMK13, produced a martian spectral analog similar to natural palagonites. The alteration products of the synthetic sample consisted of smectite, zeolite, and Fe oxides. Fe oxides were poorly
crystalline and were not completely extracted by DCB. Acidified ammonium oxalate is used to extract poorly crystalline Fe oxides in soils, while citrate-bicarbonate-dithionite (DCB) is used to extract more crystalline Fe oxides [2]. These reagents work well in soils where particles can be easily disaggregated. Palagonites, however, are not completely weathered and some of the products are still embedded in the silica-rich matrix, which prevents access to oxalate or DCB. Structure of a typical palagonite particle consists of a thin outer rim (1–5 μm) of a loose-textured mineral assemblage consisting of smectite, zeolite, and fine Fe oxides. Fe oxide in this layer can be removed using DCB. Underneath this layer is a thin, leached glass layer, where hydration and polymerization of silica and oxidation of Fe(II) take place. This layer contains nanometer-sized Fe oxide particles in various stages of formation. Although the extensively hydrated glass consists of submicroscopic pores in this layer, oxalate and DCB reagents cannot react with the Fe oxides still embedded in the silica-rich matrix. The reflectance spectra that are surface dependent change after the DCB treatment, indicating the removal of Fe(III) oxides from the surface layer. The extractants oxalate and DCB are useful in detecting the surface Fe(III) oxides and understanding the weathering process. It is very likely that the weathering on Mars is similar to that of palagonitic martian analogues, although there can be several mechanisms to yield materials with martian spectral characteristics.


AQUEOUS GÉOCHEMISTRY ON MARS: POSSIBLE CLUES FROM SALTS AND CLAYS IN SNC METEORITES. James L. Gooding, Code SN 2, Office of the Curator, NASA Johnson Space Center, Houston TX 77058-3696, USA.

Salts and Clays in SNCs: All subgroups of the shergottite, nakhlite, and chassigny (SNC) meteorites contain traces of water-precipitated minerals that include various combinations of carbonates, sulfates, halides, ferric oxides, and aluminosilicate clays of preterrestrial origin (Table 1) [1–8]. Oxygen three-isotope analysis of thermally extracted bulk water has confirmed that at least some of the water in SNCs is, indeed, extraterrestrial [9]. A mixture of aqueous precipitates found in the SNCs, comprising smectite, illite, and gypsum (with minor halite ± calcite and hematite), provides a self-consistent, though not unique, model for the bulk elemental composition of surface sediments at the Viking Lander sites (Fig. 1). Therefore, if the salts and clays in SNCs are truly linked to aqueous alteration and soil formation on Mars, then the suite of SNC secondary minerals might provide the best currently available insight into near-surface martian chemistry.

Implications for Water-based Chemistry on Mars: It is not clear whether the aqueous precipitates in SNCs represent products of surface weathering (i.e., actual rock-atmosphere reactions) or subsurface alteration (i.e., pedogenic weathering, groundwater infiltration, or deuteric alteration by residual magmatic volatiles); contributions by different water-rich environments cannot be excluded. As lavas, shergottites are the SNCs most likely to have experienced surface weathering; as deeper-seated rocks, nakhlites and chassignites may be the best candidates for deuteric or groundwater interactions. Further detailed microchemical, microstructural, and isotopic studies are needed to decipher the precise conditions under which the SNC salts and clays formed, but a few first-order conclusions about paragenesis are already possible. These inferences bear on possible geochemical reaction pathways for hydrothermal alteration, surface weathering, and soil-forming processes on Mars.

First, precipitation of the SNC secondary minerals occurred from saline liquid water. The vein-forming textures of the various hydrous and water-soluble phases are unmistakable evidence for aqueous solutions. Furthermore, on chemical-thermodynamic grounds, liquid water would be required to stabilize Fe-rich and Al-poor, smectite-type clay minerals under martian surface conditions [10]. Persistent traces of Cl and P (probably as PO43-) in Nakhlite clays and Chassigny carbonates show that the solutions contained significant concentrations of dissolved anions.

Second, the solutions were oxidizing and alkaline. Among the SNC secondary minerals, carbonate and sulfate are the dominant forms of carbon and sulfur, respectively, and much of the total iron is in the ferric state. In terms of aqueous geochemistry, the calcite-gypsum-hematite assemblage in Nakhlite (with no evidence for secondary sulfides) implies Eh > -0.2 and pH > 8 [11, p. 396]. Smectites also require pH > 6 for precipitation [12] although such conditions can be achieved by the reaction of acidic hydrothermal solutions with silicate minerals, as occurs on Earth where midocean hot springs encounter sea floor sediments [13]. Pure water in equilibrium with the Mars atmosphere should definitely be acidic so that formation of an alkaline solution on Mars would require either protracted disequilibrium with the atmosphere or extensive hydrolysis of silicates [10]; the latter process dominates chemical weathering on Earth [14].

The temperatures and pressures of the solutions from which the SNC clays and salts precipitated are not well constrained by current knowledge except that extreme hydrothermal conditions are probably excluded. If, as indicated in Nakhlite, the original Ca-sulfate among the SNC precipitates was gypsum rather than anhydrite, arguments based on equilibrium relationships among water-precipitated phases in the CaSO4-H2O system [15] would argue for formational temperatures <70°C. Although anhydrite is stable to much higher temperatures compared with gypsum, occurrence of anhydrite as the principal Ca-sulfate would not necessarily require extraordinarily high formational temperatures. Anhydrite can be precipitated directly from terrestrial seawater, for example, at tem-

| TABLE 1. Products of aqueous alteration in SNC meteorites. |
|----------------------|----------------|----------------|
|                       | Shergottite | Nakhlite | Chassigny |
| CaSO4                | [1]         | [3]      | [5,6]     |
| Mg2-bearing CaCO3    | [8]         |          |           |
| MgCO3                | [6]         |          |           |
| (Fe, Mn)CO3          | [7]         |          |           |
| CaSO4 · nH2O         | [1]         | [3]      | [5,6]     |
| (Na, K)2(SO4)2 · nH2O| [3]         |          |           |
| (Na, K)Cl            | [3,4]       |          |           |
| "illite" (K, Na)2Ca3O[(Al, Mg, Fe)5] · (Si, Al)4 · O12[SiO4]2 · H2O | [2] |          |           |
| Smectite             | [3,4]       |          |           |
| (Na, Ca)2[Al(OH)3]2 · (Al, Mg, Fe) · (Si, Al)4 · O12[(OH)2]2 · nH2O | [4] |          |           |

* x, y, and n are generalized coefficients where stoichiometry is variable or uncertain.
Fig. 1. Bulk elemental composition of martian surface sediments at the Viking Lander 1 site (open circle [20]) compared with whole-rock compositions of representative SNC meteorites (filled triangles), individual clay found in SNCs (filled circles and curved mixing line), and compositional fields of common aluminosilicate minerals. Note that the VL-1 composition should be expected to plot higher along the y-axis if its Na and K concentrations were known (i.e., Viking did not measure Na and set only an upper limit for K). By adding Na or K, the VL-1 point can be moved vertically upward onto the mixing line defined by the SNC clays; by subtracting Ca as CaSO4, it can be moved vertically downward through the smectite field. AMP = amphibole; FSP = feldspar; OL = olivine; PX = pyroxene; ZEOL = zeolite.

Figures of only about 150°C [16]. In addition, stabilization of ferric hydroxide requires temperatures <100°C [17]. At present, no lower limit can be set for the temperature at which the SNC aqueous precipitates formed and solutions at or near freezing cannot be excluded. The HO enrichment in calcite from EETA79001,239 is most reasonably attributed to reactions within the CO2-H2O system at low temperatures (where isotopic fractionations are greatest), perhaps as low as 0°C [18]. Because carbonates in EETA79001,239 have generally the same isotopic compositions as those in Nakhl, a low-temperature origin is also implied for the Nakhl carbonates. If the solutions were cold, however, the moderate crystallinity of the clays would suggest that the fluids were long-lived in order to explain the partial ripening of colloidal gels in ordered phyllosilicate phases. Ordinarily, ripening times of at least a few weeks are required to experimentally synthesize clay minerals at <200°C [19, p. 479-490]. Better thermometric constraints will require more detailed mineralogical studies and, ideally, separate oxygen-isotopic analyses of coexisting precipitates.

If the SNC parent planet is Mars, as previously inferred from independent evidence, the aqueous precipitates indicate that oxidizing, water-based solutions probably have been chemically active on Mars for at least the time interval represented by the radiometric ages of the meteorites, namely, the past 200-1300 m.y.


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EFFECT OF PURITY ON ADSORPTION CAPACITIES OF A MARS-LIKE CLAY MINERAL AT DIFFERENT Pressures.

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Introduction: There has been considerable interest in adsorption of carbon dioxide on Marslike clay minerals. Some estimates of the carbon dioxide reservoir capacity of the Martian regolith were calculated from the amount of carbon dioxide adsorbed on the iron-rich smectite nontronite under Martian conditions [1,2]. The adsorption capacity of pure nontronite could place upper limits on the regolith carbon dioxide reservoir, both present martian atmospheric pressure and at the postulated higher pressures required to permit liquid water on the surface. In this study, adsorption of carbon dioxide on a Clay Mineral Society standard containing nontronite was studied over a wide range of pressures in the absence of water. Similar experiments were conducted on the pure nontronite extracted from the natural sample. Heating curves were obtained to help characterize and determine the purity of the clay sample.

Experimental Procedure: The Clay Mineral Society standard used was SWA-1, a ferruginous smectite from Grant County, Washington. Except for routine crushing, the unpurified samples were not treated prior to use.

The natural sample was crushed and suspended in water via disaggregation by sonication. Particles <2 µm were isolated by filtration and centrifugation. The final SWA-1 product was flocculated with MgCl2. Excess salt was removed from the clay using an HCl wash, and the clay was dried in a vacuum desiccator at room temperature. The clay was ground in a porcelain mortar and pestle, and a known mass of clay was dried in a vacuum desiccator at 105°C for 24 hours. The clay was then weighed and ground in a porcelain mortar to ensure a uniform clay mass. The clay was then heated to 300°C at a rate of 1°C/minute to remove the water of hydration, and the weight of the clay was measured to determine the amount of water adsorbed at the various temperatures.

Fig. 1. Heat flow curves for natural and purified SWA-1.
Fig. 2. Comparison of carbon dioxide adsorption capacity for natural vs. purified samples.

dialysis. Water was removed by freeze drying. The resulting sample was a fluffy yellow solid. For the adsorption studies, the samples were placed in a Waring blender on the low power setting for 15 s in order to provide uniformity in the samples. Sample sizes of 0.25 g were used in each trial.

The apparatus used in the adsorption studies consisted of a stainless steel vacuum rack with a removable glass sample holder. The volume of the entire system was 90.0 cm$^3$, and the volume of the sample flask was 60.5 cm$^3$. The volume occupied by the sample was considered in the calculation of gas adsorbed.

A sample was placed in the flask, and the entire system was evacuated to a pressure of about 10 µm. To remove any residual gases, the stainless steel line was heated with a heat gun and the sample flask was warmed to 343 K (70°C) in a heating mantle during evacuation. When a constant pressure of about 10 µm was obtained, the sample chamber was brought to the desired temperature, and the valve to the vacuum pump and the stopcock to the sample flask were both closed. The sampling line (of predetermined volume) was then filled with carbon dioxide, and the pressure was measured with an electronic manometer. The stopcock to the sample flask was slowly opened, the final pressure was measured, and the amount of carbon dioxide adsorbed by the sample was calculated.

Heating curves were obtained with a Perkin-Elmer DSC-7 Differential Scanning Calorimeter (DSC) using a scan rate of 20 K/min under argon gas purge of 18.5 cm$^3$/min.

Results and Discussion: In the purification step, the yield of pure product was 36% by weight. Nontronite content of the natural clay, measured by differential scanning calorimetry, was 30% (Fig. 1). With increasing temperature, changes in heat flow indicate the release of adsorbed and interlayer water under 500 K and release of structural (hydroxy) water between 600 and 900 K.

The adsorption capacity for the purified sample was significantly greater than for the natural smectite. Figure 2 shows carbon dioxide adsorption for natural and purified SWa-1 samples. Adsorption was three times greater on the purified sample. As expected, adsorption increased with increasing pressure for all samples studied.

Conclusion: Significantly greater carbon dioxide adsorption occurred on the purified sample. Thus, when using terrestrial clay samples as analogs for martian soils, especially when making comparisons between studies, the purity of the clay mineral sample should be considered.

Studies at lower temperatures and on an additional CMS standard (NG1) are in progress. Although these studies were initiated to determine head-space pressures in sealed containers of martian soils upon heating, the adsorption data may also be useful in estimating the magnitude of the regolith carbon dioxide reservoir.

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LOW-TEMPERATURE FORMATION OF MAGNETIC IRON OXIDES. Chr. Bender Koch$^1$ and M. B. Madsen$^2$. Laboratory of Applied Physics, Technical University of Denmark, DK-2800 Lyngby, Denmark, $^2$Physics Laboratory, H. C. Ørsted Institute, Universitetsparken 5, DK-2100 Copenhagen Ø, Denmark.

Elemental analysis and magnetic measurements of the surface of Mars have indicated the presence of an iron oxide with a considerable magnetic moment [1]. Identification of the oxide phase(s) is an important subject as this may be used to identify the process of weathering on the martian surface as well as the composition of the Mars regolith itself. Consequently, we have been interested in evidence of new formation of strongly magnetic phases (e.g., magnetic, magnetite, feroxyhite) in terrestrially derived Mars sample analogues. Within the group of Mars sample analogues derived from low-temperature weathering of basalts in Arctic regions we have never observed evidence of magnetic oxides formed at the outermost weathering rind. However, in one instance where we investigated the weathering products accumulating in a crack of a basaltic stone we have found evidence of magnetite. Below we present the experimental details.

Fig. 1. Mössbauer spectra of the magnetic fraction obtained at 298 K (a, b) and 368 K (c). Spectrum B was obtained in a magnetic field of 0.6 T applied perpendicular to the γ rays. I, II, and III: see text.
The sample was collected as a loose powder from a crack in a basaltic stone that was exposed upon splitting the stone. A magnetic fraction was obtained using a ferrite magnet and studied by Mössbauer spectroscopy. All the oxide components found in the magnetic fraction were also observed in the spectra of the nonfractionated powder; however, here the relative intensity was slightly lower.

Figure 1 shows the Mössbauer spectra of the magnetic fraction obtained using different experimental conditions. The room-temperature spectra may be considered to consist of three magnetically split sextets and one ferric doublet. Roman I, II, and III denote the three sextets. The strong polarizing effect of the magnetic field on components I and II indicates the presence of at least one strongly magnetic (ferrimagnetic) oxide in the sample. Component III is due to goethite and in order to minimize overlap of component III, particularly with component II, we have obtained a spectrum at a temperature close to the Néel temperature of goethite (see Fig. 1c).

Table 1 gives the parameters of components I and II at 368 K.

The Mössbauer parameters of components I and II are similar to those of magnetite (Fe₃O₄) although they do not exactly match literature values for pure stoichiometric magnetite [2].

This study has demonstrated that magnetic iron oxide (nonstoichiometric magnetite) has formed during low-temperature terrestrial weathering. The details of the formation of magnetite is not known.

In relation to conditions of weathering on Mars it is notable that the formation of the magnetite has included precipitation from solution in a restricted microenvironment (crack) and not on the exposed surface of the rock.

Acknowledgments: Financial support from The Danish National Science Research Council and Technical University of Denmark is gratefully acknowledged.


**TABLE 1. Mössbauer parameters of components I and II at 368 K.**

<table>
<thead>
<tr>
<th>Component</th>
<th>B_m (T)</th>
<th>σ (mm/s)</th>
<th>ζ (mm/s)</th>
<th>A_m (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>47.4 ± 0.5</td>
<td>0.23 ± 0.05</td>
<td>-0.05 ± 0.10</td>
<td>26 ± 2</td>
</tr>
<tr>
<td>II</td>
<td>45.2 ± 0.5</td>
<td>0.71 ± 0.05</td>
<td>0.08 ± 0.10</td>
<td>14 ± 3</td>
</tr>
</tbody>
</table>

PRELIMINARY RESULTS OF AN EXPERIMENTAL STUDY OF THE INTERACTIONS OF BASALT GLASS AND A WATER VAPOR ATMOSPHERE: IMPLICATIONS FOR WEATHERING ON MARS. J. J. Mazer, J. K. Bates, and C. R. Bradley, Chemical Technology Division, Argonne National Laboratory, Argonne IL 60439, USA.

Models of weathering processes on the surface of Mars involve hydrothermal alteration as the primary mechanism responsible for clay formation. Previous experimental studies of basalt glass interactions with water under hydrothermal conditions (i.e., high dilution; glass-surface-area-to-solution-volume ratio (SA/V) is small) demonstrate that phyllosilicates and zeolites are the primary alteration minerals [1–3]. Gas-solid weathering is thought to be less thermodynamically favorable and relatively unimportant [4,5]; however, the experimental alteration of basalt glass under vapor hydration conditions (large SA/V) can result in the formation of clay minerals, zeolites, and hydrated calcium silicates [1].

We have undertaken a study of the reacted layers formed on basalt glasses experimentally altered under vapor hydration conditions to resolve this issue. High SA/V ratios promote reaction product buildup in solution and promote alteration mineral formation. It has previously been shown that these reaction conditions promote weathering processes similar to those found in nature for tektite glasses [6], rhyolitic glasses [7,8], and basalt glasses [1]. The hydration of basalt glass is described for 100% relative humidity experiments at temperatures at 150°C, 175°C, and 200°C for up to 400 days. Preliminary characterization of the alteration layers with analytical electron and scanning microscopy suggest that the reaction mechanism includes precipitation of a smectite clay on the outermost surface of the glass. Between the clay and the unreacted glass is an amorphous gel-like phase (palagonite). Our results provide an experimental basis for formation of 15N. In view of the relatively high O/N ratio in the martian atmosphere, relatively high 15N/N ratio are expected to arise from cosmogenic reactions. The added bonus comes from the fact that an appreciable production of 15N is expected in the martian regolith because of its thin atmosphere.

We show from model calculations that if accurate isotopic composition data become available for N and Ne, it should be possible to put robust constraints on the time averaging martian oxygen surface pressure and subsurface water in the upper layers. Cosmogenic N and Ne isotopes formed in the martian rocks would not be expected to be released to the atmosphere; however, those formed in surface and subsurface water reservoirs, as well as in any amorphous weathering products, would be released to the atmosphere. An interesting fact is that whereas the atmospheric 15N production is linear with the surface pressure of oxygen, the regolith production of 15N is not, due to atmospheric absorption of cosmic ray energy. Furthermore, cosmogenic Ne isotopes are expected to be produced primarily in the regolith. Hence, it is possible, given accurate isotopic data on N and Ne isotopic compositions in the martian atmosphere, to delineate possible temporal changes in the martian oxygen surface pressure and its regolith. Should a martian surface sample become available, the conclusions can be made very robust.

We present model calculations of expected N and Ne isotopic compositions in the martian atmosphere for a number of hypothetical models, based on scenarios for the martian regolith from Viking Lander experiments, together with Earth-based observations of Mars and studies of SNC meteorites.

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**N93-31947**
proposing that basalt glass interactions with vapor atmospheres can be an important source of clay minerals. Therefore, the process of weathering igneous rocks on the surface of Mars by contact with the atmosphere may be more important than previously thought.

Extrapolating the results of experiments performed at high temperatures (>100°C) to conditions relevant to the surface of Mars is only meaningful if the glass weathering processes are the same in both systems. Assuming identical processes are occurring on the surface of Mars and in our experiments, direct extrapolation of the results to 0°C results in an alteration rate of approximately 0.05 µm/yr. A significant effect of relative humidity on alteration rates in vapor hydration experiments has been demonstrated for rhodolitic glasses [9]. Thus, the limited amount of water in the martian atmosphere probably means that the actual weathering rate may be significantly lower.


MARS BRINE FORMATION EXPERIMENT. Jeffrey M. Moore1, Mark A. Bullock2, and Carol R. Stoker1, 1Space Sciences Division, NASA Ames Research Center, Moffett Field CA 94035, USA, 2Department of Astrophysical, Planetary, and Atmospheric Sciences, University of Colorado, Boulder CO 80309, USA.

Evaporites, particularly carbonates, nitrates, and sulfates, may be major sinks of volatiles (e.g., CO2, N2) scavenged from the martian atmosphere [1–3]. Mars is thought to have once had a denser, warmer atmosphere that permitted the presence of liquid surface water. The conversion of atmospheric CO2 into carbonate is hypothesized to have degraded the martian climate to its present state of a generally subfreezing, desiccated desert (see review in [4]). The rate for such a conversion under martian conditions is poorly known, so the timescale of climate degradation by this process cannot be easily evaluated. If some models are correct [e.g., 1,2,5,6], carbonate formation may have been fast at geological timescales. The experiments of Booth and Kieffer [7] also imply fast (10-100 yr) removal of the missing CO2 inventory, estimated to be 1–5 bar [2], by means of carbonate formation. The timing of formation of many of the fluvial features observed on Mars is, in large part, dependent on when and how fast the atmosphere changed. A knowledge of the rate at which carbonates and nitrates formed is also essential to assessing the probability that life, or its chemical precursors, could have developed on Mars [4]. No previous experiments have quantitatively evaluated the rate of solution for a suite of mobile anions and cations from unaltered minerals and atmospheric gases into liquid water under Mars-like conditions. Such experiments are the focus of this task.

The presence of water-soluble cations and anions in the martian regolith has been the subject of speculation for some time [e.g., 8]. Viking lander data provided evidence for salt-cemented crusts on the martian surface. This evidence is in the form of lander imagery of friable soil layers and planar fragments of disturbed soil [e.g., 9,10], and the detection of abundant sulfur and some evidence for chlorine in the soil [e.g., 11]. Martian sulfur is very likely to exist in the form of sulfate, and chlorine in the form of chlorides [11], chlorates, or perchlorates [12]. Following the Viking landings, there was considerable discussion of other salt-forming materials that "should" exist on the martian surface, including carbonates [e.g., 13] and nitrates [e.g., 3,14]. Very recently, Earth-based IR spectroscopic evidence has been obtained indicating the presence of carbonates as well as sulfates and other hydrates on the martian surface [15]. Carbonates have been detected in the SNC meteorites that are commonly believed to have a martian origin [16]. If the crusts observed at the two Viking landing sites are, in fact, cemented by salts, and these crusts are globally widespread, as ARTM-derived thermal inertia studies of the martian surface seem to suggest [e.g., 17], then evaporite deposits, probably at least in part derived from brines, are a major component of the martian regolith. The composition of liquid brines in the subsurface, which not only may be major agents of physical weathering [8] but may also presently constitute a major deep subsurface liquid reservoir [18], is currently unconstrained by experimental work.

A knowledge of the chemical identity and rate of production of martian brines is a critical first-order step toward understanding the nature of both these fluids and their precipitated evaporites. Laboratory experiments are being conducted to determine the identity and production rate of water-soluble ions that form in initially pure liquid water in contact with Mars-mixtures gases and unaltered Mars-analog minerals. The main components of the experiment apparatus consist of 10 identical, hermetically scalable sample containers made of Teflon. The interior volume of the containers is 250 ml. The lids of each container have two ports to which are fixed stopcocks to provide gas flow access to the head space when needed. Above one of the stopcocks a stainless steel vacuum/gas line fitting was mounted. Each of the containers will be kept within a large (and heavy) bell jar. The bell jar's mouth is sealed to its base plate with vacuum grease. The bell jar base contains two gas ports. Each port has a stainless steel gas line running from the exterior of the base. One line is attached to a cylinder of gas with the same mixture as that of the martian atmosphere. The entire assembly of bell jar and enclosed sample containers set within a large refrigerator whose interior can be maintained at 20° ± 0.5°C. The other two sample containers also reside within the refrigerator.

The experiment pilot run set consists of 10 samples. Individual pristine minerals to be simulated martian rocks were chosen on the basis of SNC meteorite mineralogy. Six of the samples are composed of 50 g of sorted and mixed minerals (57.2% augite, 25.26% forsterite, 13.14% anorthoclase, 3.14% ilmenite, 0.78% pyrite, and 0.48% chloropatite) acting as unaltered Mars rock analogs immersed in 100 ml of deoxygenated, doubly distilled liquid H2O, and above which the head space is filled with a martian gas mixture (CO2, 95.50%, N2, 2.70%, Ar, 1.60%, O2, 0.13%, CO 0.07%). These six containers will be the source of the principle data to be obtained by this pilot run set. Two other sample containers are filled with the same materials as those just discussed but with terrestrial gas in their head space. The results of the analysis of their fluids will provide some ability to compare the "martian" vs. terrestrial environmental influence, which in this case is only the difference between the two atmospheric compositions. Two other sample containers hold 100 ml of deoxygenated, doubly distilled H2O and martian gas mixture in the head space (nor rock particles). These serve as controls. The rock/gas/water or gas/water mixtures are allowed to interact with one another for specific time durations, then the reactions are stopped by the removal of the fluid, which is stored under argon in the transfer containers. The individual run durations
The eight Mars gas-containing containers were placed within the bell jar. The bell jar is flushed with Mars gas. The bell jar and the two containers with Earth gas in their head space reside within a refrigerator, whose temperature is maintained at a constant 20°C ± 0.5°C. The temperature is periodically monitored with an analog (Hg-containing) thermometer that sits within a depression on the bell jar base (exterior to the bell jar). The gas pressure within the sample containers is 1 bar.

When sample interaction is terminated, the container is removed from the bell jar. The fluids of the samples are analyzed for their pH and for a number of cations and anions. The cations to be searched for are Fe, Mg, Al, K, Na, Ca, Ti, Mn, and Ba with minimum quantity detection limits of 1 ppm. The anions to be examined are Cl, F, SO₄²⁻, NO₃⁻, NO₂⁻, and CO₃²⁻ with minimum detection limits of 10 ppm.

The results of the pilot run set will provide the first experimental determination of the rates of formation and relative abundances of the common water-soluble cations and anions that form in liquid water in contact with initially unaltered "martian" rocks and the gases of the present martian atmosphere for up to two years after initial contact. If an equilibrium composition is recognized or can be extrapolated from trends in the production rates, then a model martian brine composition will be established. Such a brine can then be synthesized in any lab. The dehydration of the model brine could be used to create a model martian evaporite.

The presence of brines on Mars has been hypothesized, both as a crustal store of H₂O-rich liquid and as an agent of chemical rock weathering. The physical properties of martian brine can be evaluated by making solutions of brine at various concentrations, guided by the results of this experiment, and then measuring such parameters as density and freezing point. The model martian brine can also be used in rock weathering experiments or calculations. An extension of this work would be to investigate the effect of brine evaporite composition on clay mineral chemistry. The spectral signature of the model evaporite can be compared with spectral data to be obtained from past, present, or forthcoming missions and observations.


Introduction: The study of palagonitic soils is an active area of research in martian geoscience because the spectral and magnetic properties of a subset are spectral and/or magnetic analogues of martian bright regions [e.g., 1-6]. An understanding of the composition, distribution, and mineralogy of ferric-bearing phases for palagonitic soils forms, through spectral and magnetic data, a basis for inferring the nature of ferric-bearing phases on Mars. Progress has been made in this area, but the dataset is incomplete, especially with respect to spectral mineralogy of palagonitic soils.

Sample and Methods: The <1-mm size fraction (obtained by wet sieving with Freon) of bulk PN-9 soil was used for this study. A citrate-dithionite-bicarbonate (DCB) method was used to extract free ferric oxides as described by [7]. Untreated and DCB-extracted samples of PN-9 were studied by diffuse reflectance spectroscopy, Mössbauer spectroscopy, and scanning electron microscopy.

Mössbauer Mineralogy: Mössbauer spectra (293 K) of PN-9 before and after DCB extraction are shown in Fig. 1. Both spectra are characterized by a ferrous doublet with quadrupole splitting (QS) of ~3.0 mm/s, a weak ferrous doublet with QS of ~2.1 mm/s, a ferric doublet with QS of ~0.70 mm/s, and a magnetic sextet with a hyperfine field (Bhf) of ~51.0 T and QS of ~0.17 mm/s. The ferric doublet with QS ~3.0 mm/s most likely results from weathered olivine. The assignment for the other ferrous doublet is pyroxene and/or glass.

Of the common ferric oxide/oxyhydroxide phases, hematite (Hm) is the most reasonable assignment for the sextet. The values of Bhf and QS for PN-9 are in good agreement with the corresponding values of 51.2 T and ~0.22 mm/s for bulk-Hm [e.g., 8]. Maghemite, goethite, and lepidocrocite are not reasonable assignments because at 293 K their bulk forms are characterized by QS = 0.0 mm/s, Bhf = 38.0 T, and Bhf = 0.0 T, respectively [e.g., 8]. The skewed shape of the lines could result from a second magnetic phase, a range in the composition of the hematite resulting from Ti substitution, and/or hematite particles in the diameter range ~20-140 nm.

The ferric doublet most likely results from nanophase ferric oxide particles (np-Ox). NP-Hm (hematite particles having diameters less than ~10 nm) is characterized by the same Mössbauer parameters [9], but unique assignment to this or any specific np-Ox is not possible.
They found Fe-rich regions in palagonitic glasses. The spectra of PN-9 before (Fig. 1a) show the glassy basalt matrix of palagonitic soil particles. The spectrum after DCB extraction (Fig. 1b) shows the procedure preferentially removed Fe-rich surface regions. The Mössbauer data identify the pigmentary phase as predominantly np-Ox particles. The position of the band at 850–900 nm and Mössbauer data are both consistent with the effect of DCB extraction on the reflectivity spectra of PN-9 is shown in Fig. 2. The untreated PN-9 sample shows a ferric absorption edge through the visible that gives the sample its characteristic brown color. There is a pronounced inflection near 480 nm, a relative reflectivity maximum near 750 nm, and a shallow band near 850–900 nm. After DCB extraction, the sample is grey because the ferric absorption edge is at shorter wavelengths. The spectral slope in the near-IR is negative and the 750-nm feature is the reflectivity maximum. A shallow band centered near 900 nm is also present. The reflectivity data clearly establish that the pigmentary phase for PN-9 is DCB extractable. The Mössbauer data identify the pigmentary phase as predominantly np-Ox particles. The position of the band at 850–900 nm and Mössbauer data are both consistent with subtraction of spectrum 1b from 1a under the constraint that the resulting spectrum has no positive peaks. Note that the ferric doublet resulting from np-Ox is present. The presence of a residual ferric doublet in the DCB-extracted sample implies some of the np-Ox was protected from the reagent, probably by being imbedded within a glass matrix as observed by [7]. The np-Ox extracted by DCB was probably derived from porous surface rinds where they are readily accessible to the reagent. This view is supported by SEM results that show that DCB treatment removed Fe-rich surface regions. Chemical analysis showed ~38% of the total Fe was removed by DCB extraction.

The effect of DCB extraction on the reflectivity spectra of PN-9 is shown in Fig. 2. The untreated PN-9 sample shows a ferric absorption edge through the visible that gives the sample its characteristic brown color. There is a pronounced inflection near 480 nm, a relative reflectivity maximum near 750 nm, and a shallow band near 850–900 nm. After DCB extraction, the sample is grey because the ferric absorption edge is at shorter wavelengths. The spectral slope in the near-IR is negative and the 750-nm feature is the reflectivity maximum. A shallow band centered near 900 nm is also present. The reflectivity data clearly establish that the pigmentary phase for PN-9 is DCB extractable. The Mössbauer data identify the pigmentary phase as predominantly np-Ox particles. The position of the band at 850–900 nm and Mössbauer data are both consistent with subtraction of spectrum 1b from 1a under the constraint that the resulting spectrum has no positive peaks. Note that the ferric doublet resulting from np-Ox is present. The presence of a residual ferric doublet in the DCB-extracted sample implies some of the np-Ox was protected from the reagent, probably by being imbedded within a glass matrix as observed by [7]. The np-Ox extracted by DCB was probably derived from porous surface rinds where they are readily accessible to the reagent. This view is supported by SEM results that show that DCB treatment removed Fe-rich surface regions. Chemical analysis showed ~38% of the total Fe was removed by DCB extraction.

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Fig. 1. Mössbauer spectra (293 K) for the <1-mm size fraction of PN-9.

Based on Mössbauer data. Direct observation of np-Ox particles in palagonitic soils has been made by [7] using transmission electron microscopy. They found Fe-rich regions ~10 nm in diameter within the glass matrix of palagonitic soil particles that were obtained from a sample whose Mössbauer spectrum was dominated by a ferric doublet.

Effect of DCB Extraction: Comparison of the Mössbauer spectra of PN-9 before (Fig. 1a) and after (Fig. 1b) DCB extraction shows the procedure preferentially removed the np-Ox. This is shown directly in Fig. 1c, which is the spectrum obtained by
the presence of well-crystalline (bulk) hematite in PN-9. However, evidence for bulk-Hm is not indicated in the reflectivity data after DCB extraction even though Mössbauer data show it is still present. A possible explanation is that the bulk Hm remaining after DCB extraction is located in particle interiors and is thus observable by Mössbauer spectroscopy but not reflectance spectroscopy.

Summary: The correspondence in spectral properties between palagonitic soil PN-9 and martian bright regions implies the nature of the pigmenting phases also corresponds. Mössbauer and reflectivity data obtained before and after DCB treatment of palagonitic soil PN-9 show that DCB removed the pigmentary phases and identify them as ferric np-Ox particles together with minor bulk Hm. The presence of bulk Hm on Mars has been inferred in other studies [e.g., 6,9-11]. The main contribution of this study and that of [7] is identification of np-Ox particles as a pigmenting agent for palagonites and, by inference, for Mars.


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SYSTEMATIC VARIATIONS IN THE SPECTRAL PROPERTIES OF BRIGHT REGIONS ON MARS. Scott Murchie, John Mustard, Janice Bishop, James Head, Carle Pieters, and Stephane Erard, Brown University, Providence RI 02912, USA, University of Paris, Orsay, France.

Introduction: Color and albedo of the martian surface define two basic surface units, dark gray material interpreted as relatively unaltered “rock” and bright reddish material interpreted as weathered “soil” [1,2]. Understanding the processes contributing to soil formation first requires assessment of the soil’s composition and compositional diversity. In this abstract, we report first results of an investigation of the character and variability of Fe- and water-bearing phases in bright reddish materials using ISM data. Based on previous information on martian surface chemistry, we also explore implications of these results for chemical evolution of martian soil.

Information on the composition and distribution of bright reddish material comes from three major sources: Viking images, measurements by the XRF and GCMS instruments on the Viking Landers, and spectroscopic data. The XRF experiment found nearly identical, Fe-rich major-element compositions comparable to weathered basalt [3]. Soil water, amounting to ~1-3 wt% as measured by the GCMS, was liberated mostly by heating to ≥350°C, suggesting that it is present in a chemically bound form [4]. Spectroscopic studies have detected ferric oxide, probably hematite [1,5,6], as well as molecular water [7]; However, the identities of major silicate phases have been controversial, with conflicting evidence regarding phyllosilicates [8,9]. Two main interpretations of this evidence have been proposed: (1) Weathering of basaltic glasses by H2O and CO2 formed a mixture of oxides, salts, and metastable phyllosilicates such as montmorillonite [10] and (2) “palagonite” formed when basaltic melt contacted ground ice or water [11,12]. Palagonite is a hydrated basaltic glass containing dispersed ferric oxide, recrystallized in varying degrees to phyllosilicates [11,13]. Typically it is aphanitic, although some samples contain phenocrysts [14]. In either case, colian redistribution is thought to have resulted in the material’s global homogenization.

Imaging spectroscopic data returned by the ISM instrument on Phobos 2 provide a powerful new basis for evaluating the composition and origin of martian soil because they are indicative of the presence and distribution of water- and Fe-bearing phases whose mineralogy is sensitive to the history of chemical weathering. ISM-returned 9 images approximately 24 x 120 pixels in size, with 22-km spatial resolution and 128 spectral channels covering the wavelength range 0.76-3.16 μm [15]. Water- and Fe-related absorptions in this range include H2O at ~1.9 and ~3.0 μm; H2O and OH at ~1.4 μm; OH at ~2.8 μm; metallic-OH at ~2.2-2.3 μm; ferric oxides and oxhydroxides near 0.9 μm; and pyroxene near 1 μm and 2 μm. Initial calibration of these data and atmosphere removal were described by Erard et al. [16]; calibrations refinements are described elsewhere [17]. The high signal-to-noise ratio within each image (up to 500:1) results in high sensitivity to the presence of weak absorptions.

Procedure: We are using Viking Orbiter images and ISM data to investigate the presence, distribution, and relation to surface geology of water- and Fe-bearing phases in bright reddish materials. Our initial investigations have covered Valles Marineris, the surrounding Tharsis plateau, and Arabia, which exhibit a variety of
Spectral Heterogeneities in Bright Reddish Materials: The strength of the 3-μm H₂O absorption exhibits lateral variations of several percent within bright reddish materials (Fig. 1). The variations are spatially coherent and independent of albedo and atmospheric path length, thus representing physical variations rather than calibration or optical artifacts. In fact, the variations are correlated with surface geology. "Normal" bright areas, with less strong absorptions, correspond to surface morphologies indicative of only a thin cover of dust and soil (e.g., cratered plains). In contrast, "hydrated" bright regions with stronger absorptions correspond to unconformable deposits in Candor Chasma and western Arabia. Both exhibit stronger 3-μm H₂O absorptions than do dark gray regions.

Representative spectra synoptically illustrate the "normal" and "hydrated" regions' spectral systematics (Fig. 2). The H₂O absorption at 1.9 μm is too weak (≤0.3–0.5%) to observe. However, there is a weak, narrow absorption at 2.2 μm with a depth of up to 1.5%, which meets the above criteria for validity. This absorption is attributed to ALOH bonds in a phyllosilicate. It is present in "normal" bright regions, but weaker or absent in the two "hydrated" regions as well as most or all dark gray regions. Within and around each "hydrated" region, strengths of the 2.2-μm and 3-μm absorptions are inversely correlated (r ≈ −0.7) (Fig. 1a). This relationship substantiates identification of the 2.2-μm absorption, and provides information on local variations in the chemical form of water: If strength of the 3-μm absorption is indicative of water content, then more water-rich bright materials are also less hydroxylated.

"Normal" bright regions exhibit absorptions consistent with one or more ferric-bearing minerals. Most have an absorption ~3–4% deep with a band minimum near -0.86 μm, consistent with hematite. Eastern Arabia is distinct, with a stronger (~5–6%) absorption having a band minimum near -0.93 μm. In contrast, "hydrated" regions' Fe absorptions are broadened toward longer wavelengths: Candor Chasma exhibits two minima, at -0.86 and -0.96 μm, and western Arabia exhibits a minimum at -0.95 μm. Both "hydrated" regions also exhibit a broad, weak absorption centered at ~2.1 μm that is weaker or absent in "normal" bright regions (Fig. 1b). These observations suggest that the "hydrated" regions also contain ferric iron plus pyroxene as an additional Fe-containing phase.

Water-related Absorptions in Laboratory Analogs: At room temperature, Fe-montmorillonites contained ~7–11% H₂O and palagonites 13–24%, with the large water contents producing nearly saturated 3-μm absorptions. All samples exhibited significant 1.4-, 1.9-, and 2.2-μm absorptions, which were stronger in the clays, presumably due to their greater crystallinity [cf. 10]. The 2.2-μm absorption was consistently the weakest water-related absorption. After heating to ~275°C, the analogues acquired more Mars-like
water contents and spectral properties. $\text{H}_2\text{O}$ content dropped to $\leq 1\%$ for the clays and 3–4\% for the palagonites. Three-micrometer band strengths decreased in all cases, but by the largest amounts in the clays (Fig. 3). Strengths of the 1.4- and 1.9-$\mu$m absorptions decreased to less than that of the 2.2-$\mu$m absorption; the clays retained a stronger 2.2-$\mu$m absorption than in the palagonites. These and similar results reported by Bruckenthal [21] are understandable in terms of crystallinities of the analogues: $\text{H}_2\text{O}$ incorporated in the glassy structure of palagonites is held more tightly than electrostatically bonded water in crystalline clay. As a result, under dehydrating conditions palagonites exhibit greater water contents and stronger 3-$\mu$m absorptions than clay. Conversely, the less crystalline palagonite exhibits a weaker 2.2-$\mu$m absorption.

Possible Compositional Interpretations: In light of Viking Lander elemental abundances and the experiments with laboratory analogues, the 2.2-$\mu$m absorption in "normal" bright reddish material is interpreted to indicate presence of dehydrated, iron-rich montmorillonite. This interpretation is supported by 0.4–1.1-$\mu$m reflectance spectra of the laboratory analogues, which demonstrate that montmorillonite saturated with ferric iron reproduces Mars’ characteristic spectrum in that wavelength range [22]. However, regional variations in the 0.9-$\mu$m Fe absorption imply a degree of chemical heterogeneity inconsistent with global homogenization.

The relationship of "normal" and "hydrated" bright materials is constrained by the stronger 3-$\mu$m absorption, weaker 2.2-$\mu$m absorption, and stronger 2-$\mu$m pyroxene absorption in the "hydrated" regions. The "hydrated" regions’ weaker 2.2-$\mu$m absorption and stronger 2-$\mu$m pyroxene absorption cannot be explained simply by mixing of dark gray and "normal" bright materials because of the weak 3-$\mu$m $\text{H}_2\text{O}$ absorption in dark gray material. However, at least two models can explain spectral differences between the two types of bright reddish material as resulting from soil-forming processes.

Model 1. "Hydrated" materials consist of "normal" bright material and basalt fragments, cemented by hydrated salts and perhaps ferric oxyhydroxides into a "duricrust" like that at the Viking Lander sites [18]. The intergranular cement masks the 2.2-$\mu$m absorption.

Model 2. "Hydrated" materials are deposits of palagonitized basaltic glass containing pyroxene phenocrysts, from which a glassy fine-grained fraction has been partially removed and redistributed regionally by wind. Reaction of the fines with $\text{H}_2\text{O}$ and CO$_2$ formed montmorillonite [cf. 10], which became dehydrated to form "normal" bright material.

We are currently expanding our studies to additional bright reddish regions, to further determine the character of spectral heterogeneities and to evaluate models for their origin.


COMPOSITION OF WEAKLY ALTERED MARTIAN CRUST: CLUES FROM IMAGING SPECTROSCOPY.

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Introduction: Two of the fundamental questions regarding chemical weathering on Mars are the chemical pathways by which pristine crustal rocks are altered to produce observed ferric iron-bearing assemblages and inferred clay silicate, sulfate, and magnetic oxide phases, and the timing of the oxidative weathering. High-spatial- and -spectral-resolution imaging spectrometer data provide a tool to investigate aspects of these questions for material exposed on the surface of Mars. In this study, ISM imaging spectrometer data [1,2] are used (1) to develop an inventory of least-altered crustal materials and (2) to quantitatively model the mineralogic composition of the weakly altered crustal rocks.

Spectral Data: High-spatial-resolution (22 km/pixel) imaging spectrometer data were acquired in 1989 by the ISM instrument on board the Phobos II spacecraft for nine different areas near the equatorial regions on Mars [1,2]. The data windows are 300 km wide and 2000 km long and sample a wide variety of terrain and surface morphologies. In particular, the geologic units in the Tharsis plateau and Valles Marineris were well sampled with additional coverage in the Arabia and the Syrtis Major-Isidis regions. A 128-channel reflectance spectrum from 0.76 to 3.16 $\mu$m was obtained for each surface element of which 64 are used in these analyses. The signal to noise ratio is >300:1 across most of the detector elements and the absolute radiometric accuracy is $\pm 10\%$. Details of data reduction and calibration are presented elsewhere [2,3].

Composition of "Fresh Crustal Materials": Studies of the reflectance properties of Mars suggest that most regions at the scale of an ISM pixel show evidence of ferric material being present, which suggests that altered materials are widespread. Small amounts of ferric oxide can dominate the visible spectral properties, but have relatively little effect at longer wavelengths. Surfaces that are weakly altered, or relatively "fresh," can be identified based on several spectral properties. First, studies of Viking lander and orbiter images demonstrate that, in general, low-albedo materials exhibit the least effects of alteration [4,5]. Second, substantial amounts of ferric oxides would be indicated by an absorption centered near 0.85 $\mu$m or a broadening of ferrous absorptions in this wavelength area. Finally, flat continuum slopes are generally thought to indicate an absence of coating of rock surfaces by ferric oxides, although a negative continuum slope does not uniquely confirm an altered surface. Initial investigations of the ISM windows have identified many spectra that meet one or more of these criteria. A subset is shown in Fig. 1.

These spectra represent a broad sample of plains-forming geologic units of different age and surface morphology and all show a strong, well-defined absorption near 1.0 $\mu$m and an absorption of variable strength and definition near 2.1 $\mu$m. In the spectra that exhibit the strongest absorptions (Nili Patera, Melas Chasma, Eos Chasma), both the 1.0- and 2.1-$\mu$m absorptions are very symmetric and they all exhibit a shoulder on the long wavelength side of the 1.0-$\mu$m absorption near 1.15 $\mu$m. The shape, position, and strength of these absorptions are typical of calcic pyroxene [6,7] and unambiguously confirm tentative identification of this mineral phase on the martian surface made by previous workers [8,9]. The other spectra show diversity in the shape, position, and relative strengths of these two dominant absorptions, suggesting additional mineral phases or compositions are present. In particular, there is great diversity in all
these spectra in the existence, position, and character of the shoulder between 1.1 and 1.3 μm. This feature can be due to many factors including Fe site populations in pyroxene and the presence of such mineral phases as feldspar, olivine, and glass.

The dominant mineral phase spectroscopically is pyroxene. Reflectance spectra of pyroxenes exhibit variations in the wavelength position of the absorption band minima centered near 1.0 and 2.0 μm that are systematic with composition [6,7,10]. This provides a framework for deriving estimates of the pyroxene chemistry for the surface of mafic-rich regions containing pyroxene. Spectra from Nili Patera on Syrtis Major have been modeled with the Modified Gaussian Model (MGM) [11] to derive quantitative absorption parameters to compare to the results of [7] and derive initial estimates of pyroxene chemistry [3,12]. An MGM solution to the spectrum from Nili Patera is shown in Fig. 2. The data appear to be well fit by a complement of bands for a single pyroxene, although there are very small, systematic errors in the fit to the 2.0-μm band that may indicate an additional pyroxene mineral is present [13]. The band minima determined from this analysis are compared to generalized band minima chemical composition diagrams of [7] from which a Ca/(Ca + Fe + Mg) ratio of 0.275 ± 0.075 and a Fe/(Fe + Ca + Mg) ratio of 0.3 ± 0.1 are derived. The general pyroxene field that these chemical ratios correlate to is shown in Fig. 3.

Discussion: The basic results determined above indicate a basaltic composition for weakly altered crustal materials sampled by ISM. The estimate of bulk pyroxene composition determined above for Nili Patera lies between the pyroxenes augite and pigeonite. Since pyroxenes of such a composition are unusual, this may indicate a mixture of augite and pigeonite, or it may indicate a chemically zoned pyroxene with a similar bulk composition. Such zoned pyroxenes are typical in lunar basalts, which also happen to have very similar spectral properties to the low-albedo regions of Syrtis Major [3]. The strongly zoned pyroxenes in lunar basalts are considered to have resulted from rapid, nonequilibrium cooling during eruption and emplacement of the basalts. A similar set of conditions is likely for volcanics on the Syrtis Major plateau and elsewhere. Evidence for mineral phases in addition to pyroxene is subtle and the focus of continuing analysis. However, our expectations are that plagioclase will be common, as well as opacite phases and perhaps olivine. We will be continuing detail mineralogic modeling of the low-albedo regions on Mars and extend the analysis performed at Nili Patera to the other regions identified above. This will quantify the diversity of weakly altered crust, and therefore the starting materials for models of chemical weathering on Mars.


Fig. 1. (a) Representative spectra of low-albedo terrains on Mars interpreted to be of weakly altered crustal materials; labels along the right side give the geographic location from where they were collected. (b) The same spectra as in (a), but with a straight-line continuum removed to emphasize the absorption features. The same symbols are used in both plots. Note the very diagnostic, broad pyroxene absorptions near 1.0 and 2.0 μm.

Fig. 2. Fit of the Modified Gaussian Model [11] to the reflectance spectrum of Nili Patera. The actual data are shown by the crosses, the fit by the solid line superimposed on the crosses, the continuum by the dashed line, the model absorptions by the solid curves near the zero reference, and the error of the fit by the solid lines near the top of the plot.

Chemical Transport during Formation and Alteration of Martian Impact and Volcanic Deposits. H.E. Newsom, Institute of Meteoritics and Department of Geology, University of New Mexico, Albuquerque NM 87131, USA.

Much of the surface of Mars, including volcanic and cratered terrains, probably experienced alteration and degassing processes. These processes may have depleted or enriched many important elements in surface materials, including bedrock, dust, and soils. The composition of the martian soil may represent the best estimate, for some elements, of the average composition of the martian crust, similar to the composition of loess created by glacial action on the Earth [1,2]. The martian soil may represent the only convenient, globally or regionally averaged sample of the martian crust. In order to understand the composition of the source material for the soil, however, we need to understand the contributions of volcanic vs. impact sources for this material and the chemical fractionations involved in its production. The processes to be addressed include (1) degassing of volcanic deposits, as observed in the Valley of Ten Thousand Smokes at Katmai, Alaska [3,4]; and degassing of melt-bearing impact ejecta as inferred for suevite ejecta sheets at the Ries Crater [5,6], and (2) alteration or palagonitization of volcanic deposits, as documented for volcanos in British Columbia [7] and many other volcanic terrains, and impact crater deposits [5,6,8]. The process of palagonitization has been the subject of several studies with reference to Mars and palagonite is a good analogue for the spectroscopic properties of the martian dust. The role of impact cratering has not been as well studied, although Allen et al. [8] and Newsom et al. [5,6] have established that both degassing and alteration are common features of impact crater deposits. Other relevant sources of experimental data include the extensive literature on the corrosion of nuclear waste glass [9] and leaching of shocked materials [10].

Chemical Transport in Martian Analogue Systems: There are at least two processes that may result in significant chemical transport in volcanic and impact deposits. The first process is the initial degassing, including the release of volatiles from molten or partially molten fragments. The second process is the release and transport of mobile elements by hydrothermal alteration.

1. Degassing of volcanic deposits and impact-melt-bearing deposits. In studies of the fumaroles at the Valley of Ten Thousand Smokes in Katmai, Alaska, Papke et al. [3,4] have demonstrated that vapor-phase transport of both major and trace elements has occurred. They have studied fumarole deposits located in proximal (dacite) and distal (rhyolite) portions of the 1912 eruption sequence. Significant enrichments are observed for SO_4, F, H_2O, Zn, As, Cu, Pb, Br, Sn, Sb, and Au with minor enrichments for Ni and Cr. In the rhyolite portions the same elements are observed to be enriched, along with Fe, Mg, Ca, Sr, V, and Sc. Although such enrichments are undoubtedly small volumetrically, some of the fractionations are so large that a general enrichment of broad surface areas of volcanic terrains on Mars are possible. For example, enrichment factors of nearly 50 or greater have been observed for F, V, Sn, Br, As, and Pb in Katmai fumaroles. Equally important is the information that many elements are not observed to be significantly fractionated by the fumarole activity in the dacite, including Si, Ti, Al, Fe, Mg, Mn, Ca, Na, K, Ba, Sr, Ga, Y, and Sc. Some minor depletions of the REE were observed in the altered dacite. In the rhyolite portion, one fumarole that was not subjected to acid alteration showed little fractionation of the REE, while another fumarole affected by acid alteration showed mobilization of all the REE.

Chemical transport within impact crater deposits has not been extensively studied. Transport of volatile elements such as K, Mn, and Br has been reported in impact melt deposits from the East Clearwater Crater [11] and at Brent Crater [12]. These studies were of impact melt deposits located within the impact crater itself, although the impact-melt-bearing ejecta is the material that will be most subject to erosion on Mars. A characteristic of the impact-melt-bearing breccia (suevite) at the Ries Crater in Southern Germany, which is analogous to some fumaroles in ignimbrites, is the presence of degassing pipes. These pipes may represent important conduits for chemical transport. Newsom et al. [5] discussed the possible origin of these pipes by fluidization, and chemical studies will also contribute to understanding the origin of these pipes. They observed the presence of vertical degassing pipes at most of the available outcrops of suevite at the Ries, but the pipes are best exposed at the Oting quarry.

2. Hydrothermal alteration and palagonitization of volcanic and impact deposits. A likely analogue for the martian soil is the alteration product of basaltic glass called palagonite [13,14]. Palag- onite is an amorphous or microcrystalline gel-like product formed by a hydrolytic dissolution/precipitation mechanism that results in the formation of well-defined surface layers [15-17]. Work on the formation of palagonite has been stimulated by the similarity of basaltic glass to the types of borosilicate glass proposed for high-level nuclear waste disposal. The rate of formation of palagonite depends on the environment, especially the amount of silica in the fluids. Palagonitization is rapid (3-20 μm/1000 yr) on the seafloor in the presence of silica-undersaturated conditions, but is 2-3 orders of magnitude slower in the silica-saturated conditions often found in drill cores [9]. The palagonite is initially amorphous, but becomes microcrystalline with age and, in the case of freshwater and shallow seawater, palagonite is commonly replaced with authigenic clays and zeolites, usually chabazite (CaAl_2SiO_5·6H_2O). Zeolites are indicators of high reaction progress and are associated with relatively thick palagonite rinds, aluminum loss during palagonitization, and closed-system (low flow rate) conditions during alteration [7].
Chemical fractionations involving major elements during the formation of palagonite have been the subject of several studies, mainly using microprobe techniques [e.g., 18–22]. In general, Si, Mg, and Ca are depleted in the palagonite, while Fe, Ti, and Al are generally unaffected. In addition, with freshwater exposure, Na and K are generally depleted, while these elements are not depleted during palagonite formation in seawater. Similar results have been obtained in experimental alteration of basaltic glasses [e.g., 17,23]. Trace-element data are available from the freshwater alteration of basaltic glasses from British Columbia [7]. The palagonites for all samples from this study are enriched relative to fresh basalt for Cs, Ba, Hf, Ta, Th, U, and REE. Other elements, including Sr, Rb, Cr, Co, and Ni, are variably enriched or depleted in different samples, with a tendency for Co, Ni, and Cr to be more highly concentrated in high-Al palagonites, and depleted in low-Al palagonites. Other studies of trace-element fractionations during palagonitization have been reported by Staudigel and Hart [18] indicating, for example, that the REE are depleted in marine palagonites, and by Furnes [24] of subglacial hylaeolasticite from Iceland, who found that the elements least affected by alteration are Zn, Ni, Y, Ba, and Nb.

Hydrothermal alteration of impact melt sheets and impact-melt-bearing ejecta may have implications for the evolution of the surface of Mars [5,6,8,25–27]. Current estimates suggest a minimum global thickness of impact melt on Mars of 110 m [28]. Evidence of hydrothermal alteration of impact melt deposits on the Earth is ubiquitous [e.g., 8,29]. At the Ries Crater, Newsom et al. [5] showed that the mineralogy of the clays in the suevite at the Otting quarry is consistent with a relatively low-temperature formation, presumably after liquid water had become stable near 100°C. This led to the problem of establishing that the alteration was connected with the thermal event. A paleomagnetic and rock magnetic study [6] provided evidence for hydrothermal alteration occurring immediately following deposition of the impact-melt-bearing breccias (suevite). They found evidence for the formation of maghemite during low-temperature oxidation of titanomagnetite. The maghemite, however, records the same magnetization direction as Ti-magnetite, which recorded the magnetic direction when the suevite was at high temperature. These observations suggest that the low-temperature alteration of the suevite, including clay formation, occurred during hydrothermal alteration shortly after emplacement of the suevite, and not in response to weathering under ambient conditions since crater formation, 15 m.y. ago. Now that the existence of low-temperature hydrothermal alteration of impact-melt-bearing deposits has been established, further work is needed to understand the chemical fractionation that occurs during the formation of the alteration phases. Ries et al. [29] report major-element analyses for the <2-μm size fraction, which can be compared with the composition of the impact melt glass from the same outcrops. They found that Al, Mg, and Fe are usually enriched in the fine-grained fraction, Ti is usually conserved, Si and Ca show minor depletions, and Na and K are usually highly depleted.

Given that the mechanisms for alteration are similar, assuming similar types of parent material, the chemical fractionations observed in impact deposits could be similar to those observed in volcanic deposits. Several aspects of impact deposits on Mars, however, may produce different results from alteration of mafic volcanic material. The alteration of experimentally shocked mineral grains has been shown to be significantly enhanced compared to unshocked mineral grains [10]. The alteration of impact melt glasses may also be enhanced due to the formation of metastable glasses [30]. In addition, impact-melt-bearing deposits on Mars may have much larger fractions of impactor material (up to 40%) compared to terrestrial impacts (8%). Therefore, if shock-activated weathering of impact-produced glass is a primary source of soils on Mars, then those weathered martian surface materials should be greatly enriched in meteoritic material, which is consistent with the Viking XRF analyses [30].

3. Chemical transport and soil formation: Volcanic processes vs. cratering processes on Mars. The martian dust may be derived from a mixture of altered material at ancient impact sites and altered volcanic material [31]. For example, Squyres et al. [32] have suggested the possibility of large-scale volcano-ground ice interactions on Mars to produce palagonite. Additional investigation of chemical fractionation and transport within impact-melt-bearing deposits is needed to find elements that may discriminate among the alteration mechanisms, including alteration under ambient conditions. From the limited data discussed above, there is little difference in the major-element fractionations observed in suevite and the formation of freshwater palagonite, with the only difference being a slight enrichment of Fe and Al in the altered impact material, which is not present in the palagonite. Compared to the freshwater alteration, however, deep-sea palagonite does not show the pronounced Na and K depletions seen in the altered suevite and the freshwater palagonite. There may also be a distinction in the transport of elements by fumarole-type activity, because transport of the volatile elements K and Mn has been reported in impact melt deposits from the East Clearwater Crater [11] and at Bregt Crater [12]. These elements, however, were not enriched in the altered fumarole material in the dacite Droyolith at Katmai.

Conclusions: Understanding the chemical processes involved in the formation of the martian soil may be critical to interpreting chemical data obtained from orbital instruments, landers, and sample return missions in order to determine the bulk composition of the planet. Chemical clues may also be important in sorting out the relative importance of volcanic vs. impact crater sources of the martian soil and the alteration mechanisms, including alteration under ambient conditions vs. hydrothermal conditions. The abundance and distribution of water on Mars is another problem that also concerns issues such as the origin of life and the density of the early martian atmosphere [32,33]. The chemical fractionations associated with hydrothermal alteration such as palagonitization of volcanic terrains or alteration of impact deposits, if they can be observed by remote sensing, could provide clues to the distribution of water on Mars.

REFLECTANCE SPECTROSCOPY AND GEOMETRY OF PALAGONITE AND IRON-RICH MONTMORILLONITE CLAY MIXTURES: IMPLICATIONS FOR THE SURFACE COMPOSITION OF MARS. J. B. Orenberg, J. Handy, and R. Quinn, Department of Chemistry and Biochemistry, San Francisco State University, San Francisco CA 94132, USA.

Because of the power of remote sensing reflectance spectroscopy in determining mineralogy, it has been used as the major method of identifying a possible mineral analogue of the martian surface. A summary of proposed martian surface compositions from reflectance spectroscopy before 1979 was presented by Singer et al. [1,2]. Since that time, iron-rich montmorillonite clay [3,4], nanocrystalline or nanophase hematite [5], and palagonite [6–8] have been suggested as Mars soil analogue materials.

Palagonite in petrological terms is best described, perhaps, as an amorphous, hydrated, ferric iron, silica gel. Montmorillonite is a member of the smectite clay group and its structure is characterized by an octahedral sheet in coordination with two tetrahedral sheets in which oxygen atoms are shared. The crystallinity of montmorillonite is well defined in contrast to palagonite where it is considered amorphous or poorly crystalline at best.

Because of the absence of the diagnostic strong 2.2-μm reflectance band characteristic of clays in the near-infrared (NIR) spectrum of Mars and palagonite, and based upon a consideration of wide wavelength coverage (0.3–50 μm), Roush et al. [9] concluded that palagonite is a more likely Mars surface analogue. In spite of the spectral agreement of palagonite and the Mars reflectance spectrum in the 2.2-μm region, palagonite shows poor correspondence with the results of the Viking LR experiment [10,11]. In contrast, iron-rich montmorillonite clays show relatively good agreement with the results of the Viking LR experiment [3,10,12].

This spectral study was undertaken to evaluate the spectral properties of mixtures of palagonite and Mars analogue iron-rich montmorillonite clay (16–18 wt% Fe as Fe₂O₃) as a Mars surface mineralogical model. Mixtures of minerals as Mars surface analogue materials have been studied before [2,8], but the mixtures were restricted to crystalline clays and iron oxides.

Reflectance spectra from 0.3 to 2.5 μm were recorded on a Perkin Elmer Lambda 9 spectrophotometer (Norwalk, CT) using a Labsphere DRTA-9A Diffuse Reflectance and Transmittance Accessory (North Sutton, NH). Reflectance data presented below thus represent hemispherical reflectance. The spectral bandpass was set between 1/5 and 1/10 of the width at half height of the spectral features of interest by setting the slits to 2.0 nm in the UV/VIS. This allowed for a constant spectral resolution (±1%) in the UV/VIS. In the NIR, an automatic slit program was used to maintain a constant energy level during spectral scanning (120 nm/min).

Mixtures (% by wt) of palagonite with the iron-rich Mars analogue montmorillonite (15.8 ± 0.8 wt% Fe as Fe₂O₃—a full Mars iron analogue) are shown in Fig. 1. In the very important 2.2-μm region, the band due to clay lattice structure becomes noticeable in mixtures at the 10–20 wt% Fe-montmorillonite level. In order to evaluate this observation more quantitatively, a rigorous band depth analysis was carried out [12]. The results indicated that band depth at 2.2 μm is insensitive to the presence of up to 15 wt% Fe-

![Fig. 1. A comparison of the diffuse reflectance spectra of different wt% mixtures of palagonite and Fe-montmorillonite (15.8 ± 0.8 wt% Fe as Fe₂O₃). All spectra are scaled to unity at 1.02 μm.](image-url)
montmorillonite. Above these concentrations, there is an increase in band depth with increasing wt% Fe-montmorillonite (decreasing palagonite), which is attributable to the 2.20-μm absorption feature characteristic of smectite clays. If one accepts the premise that palagonite is a “good” spectral analogue of the Mars surface material, up to 15 wt% of Fe-montmorillonite can be present on the surface of Mars and remain undetected. In spite of the fact that the most recent telescopic observations of Mars do not show evidence of a 2.20-μm band [13], the absence of the 2.20-μm band cannot be used to eliminate less than 15 wt% iron-enriched montmorillonite. The conclusion follows that a Mars analogue, iron-rich, montmorillonite clay can be present on the surface of Mars as a major component (up to 15 wt%) of the Mars soil even if the 2.20-μm band is absent from remotely sensed spectra.

Simulations of the Gas Exchange Experiment (GEX), one of the Viking Lander Biology experiments [14,15], were run using palagonite and iron-enriched montmorillonite. Periodic sampling of the headspace gases above the mineral samples and analysis using gas chromatography was carried out with mineral samples in dry, humid, or wet conditions using a nutrient solution of the same composition as used in the Viking Lander experiments. Interpretation of the data shows that a correspondence to the Viking results occurs with palagonite-iron-montmorillonite mixtures. The results will be discussed with reference to the conclusion of the reflectance spectroscopic measurements and previous results of the Labeled Release Viking Biology Experiment [3,10].


**USING THE VIKING BIOLOGY EXPERIMENTAL RESULTS TO OBTAIN CHEMICAL INFORMATION ABOUT MARTIAN REGOLITH.** Robert C. Plumb, Worcester Polytechnic Institute/ Brown University, Providence RI, USA.

Although initially formulated as biology experiments, most of the results produced by the Viking Labeled Release (LR), Gas Exchange (GEX), and Pyrolytic Release (PR) experiments have been reproduced by chemical means [1]. The experiments do not need more study as “biological” phenomena, but they do deserve much more careful consideration from a chemical viewpoint. They are the only “wet-chemical” experiments that scientists have performed on another planet, but they have not found very general use as sources of scientific information. There is a large set of potentially useful chemical observations, e.g., the three resolveable and precisely measured kinetic components of the release of 14C-labeled gases, the thermal sensitivity and magnitudes of the oxidation reaction(s) of the LR experiments, the kinetics and magnitude of the O2 and CO2 release of the GEX experiments, the thermal sensitivity of the GEX results, the differences between the thermal sensitivity of the GEX and the thermal sensitivity of the LR responses, and the kinetics and magnitudes of the LR successive injection reabsorption effect. It should be possible to test many chemical aspects of hypothetical martian phenomena in experiments using the biology experimental configurations and derive much valuable information by comparisons to the Viking observations.

To a considerable extent the underutilization has resulted from the fact that the methods of scientific logic tend to become tailored to specific fields of study [2]. The methods by which information can be derived from the Viking biology experiments do not match well with the currently popular logic methodologies used by chemists and other scientists: The experiments were not designed by chemists; they were biology experiments. However, the necessary methods of "chemical logic do exist.

It is suggested that the scientific community would benefit and obtain more information from the Viking results if we were to resurrect a mode of thinking that was widely used in the nineteenth century. It isn’t a standard deductive or inductive process, but is a method of reasoning specifically developed to take advantage of observations of chemical reactions. Scientists need to understand the chemical subtleties in this method if they are to effectively judge arguments that invoke the Viking biology results. A superb explanation of this chemical logic was provided by Michael Faraday in his lecture series to groups of young people ca. 1850 entitled "Chemical History of a Candle" [3]. In those lectures Faraday showed how comparisons of the chemical reactions of products formed when a candle burns in air with those reactions occurring with products formed when carbon and hydrogen are burned in air can be used to prove that the wax of a candle is composed of carbon and hydrogen. A similar logical method has been used (with decreasing frequency in recent years) in high school and college qualitative analysis courses where the similarity of precipitates and colors produced by reactions of unknowns and knowns with various reagents has been used to determine what cations and anions are present in an unknown salt. Nowadays basing a proof upon observations of chemical reactions has fallen into disuse and the power of the method is not generally appreciated. Today’s scientists do not resort to these “old-fashioned” proofs; they pride themselves on state-of-the-art apparatus. They would send such samples to the analytical lab for mass spectrometry or atomic absorption tests and draw conclusions by “deductive” arguments based on such physical measurements.

Several aspects of Faraday’s logical process must be given careful consideration in designing laboratory studies of the Viking biology results and in drawing conclusions from the experiments. Chemical reaction rates and magnitudes of reactions at equilibrium are usually very sensitive to physical parameters such as temperature, pressure, methods of product detection, and reactor vessel geometry. Unless the correct physical parameters, including the trapped atmosphere of Mars, are built into the simulations, the results can lead to erroneous conclusions [4]. Even such seemingly minor matters as the conditions of storage after sample acquisition and before testing can impact the results, it having been demonstrated that absorbed H2O2 often suggested to explain the LR and GEX results, does not survive the conditions of storage [5].

After establishing that the experimental conditions of the test satisfactorily mimic the actual Viking biology experiments one needs to consider the “resolution” of the tests, i.e., could more than one chemical model produce the same quality of simulation? The resolution, and hence the quality of proof for a particular model, increases greatly with the number of independent observations being replicated. Much of the power of this approach to determining the
CHEMICAL MODELING CONSTRAINTS ON MARTIAN SURFACE MINERALOGIES FORMED IN AN EARLY, WARM, WET CLIMATE, AND SPECCULATIONS ON THE OCCURRENCE OF PHOSPHATE MINERALS IN THE MARTIAN REGOLITH. Geoffrey S. Plumlee¹, W. Ian Ridley¹, and Jeffrey D. DeBraal² ¹U.S. Geological Survey, MS 973, Denver Federal Center, Denver CO 80225, USA, ²Department of Geological Sciences, University of Oregon, Eugene OR 97403, USA.

This is one in a series of reports summarizing our chemical modeling studies of water-rock-gas interactions at the martian surface through time. The purpose of these studies is to place constraints on possible mineralogies formed at the martian surface and to model the geochemical implications of martian surficial processes proposed by previous researchers. Plumlee and Ridley [1] summarize geochemical processes that may have occurred as a result of inferred volcanic- and impact-driven hydrothermal activity on Mars. DeBraal et al. [2,3] model the geochemical aspects of water-rock interactions and water evaporation near 0°C, as a prelude to future calculations that will model sub-0°C brine-rock-clarathrate interactions under the current martian climate.

In this report, we discuss reaction path calculations that model chemical processes that may have occurred at the martian surface in a postulated early, warm, wet climate [4]. We assume a temperature of 25°C in all our calculations. Processes we model here include (1) the reaction of rainwater under various ambient CO₂ and O₂ pressures with basaltic rocks at the martian surface, (2) the formation of acid rain by volcanic gases such as HCl and SO₂, (3) the reactions of acid rain with basaltic surficial materials, and (4) evaporation of waters resulting from rainwater-basalt interactions.

Theory of Chemical Reaction Path Modeling: Computer-based chemical reaction path calculations determine heterogeneous chemical equilibrium conditions among minerals, gases, and/or aqueous solutions along an incremental series of specified temperature, pressure, and/or bulk composition changes [5]. Based upon measured or estimated thermodynamic data, reaction path modeling predicts the changes in aqueous solution composition and the resulting amounts of minerals precipitated during specific chemical processes. Because they are computer-based, large numbers of reaction path calculations can be carried out to address the full range of factors influencing a particular chemical process. The chemical speciation and reaction path programs SOLVEQ and CHILLER [5,6] were used in this study.

It should be noted that minerals predicted by the modeling to be stable thermodynamically for a given set of chemical conditions may not actually precipitate in nature due to kinetic factors. For example, Nordstrom et al. [7] note that quartz, illites, smectites, and other aluminosilicates are not likely to precipitate at 25°C except under quite saline conditions; instead, minerals such as chaledony, kaolinite, and aluminum hydroxides are kinetically stable. Also, ferrihydrite (rather than goethite or hematite) probably controls iron solubility under oxygenated conditions. We have addressed such considerations in the modeling by carrying out both runs in which kinetically inhibited minerals were allowed to precipitate, and runs in which precipitation of these minerals was suppressed, thereby allowing the precipitation of thermodynamically less-stable minerals.

Constraints on the Geochemistry of Martian Rainwater: The initial chemistry of rainwater strongly affects the minerals formed during rock weathering. It is thus necessary to constrain the chemistry of martian rainwaters in a warm, wet climate. Lacking detailed information in this regard, we have used in our calculations a range of water compositions reflecting (1) reasonable variations in ambient partial pressures of atmospheric CO₂ and O₂ and (2) possible variations in the input of acidic volcanic gases such as SO₂ and HCl. For example, under current Earth conditions, rainwater in equilibrium with atmospheric O₂ (pO₂ = 0.2 bar) and CO₂ (pCO₂ = 0.033 bar) has calculated pH = 5.66, dissolved O₂ = 2.5 x 10⁻⁴ m/kg, and dissolved CO₂ = 1.3 x 10⁻³ m/kg. In contrast, martian rainwaters in a postulated warm, wet climate with pCO₂ = 2.2 bar [4] and pO₂ = 10⁻⁵ bar (approximately the same as that for present-day Mars) are calculated to have lower pH (4.08), greater dissolved CO₂ (7.5 x 10⁻² m/kg), and lower dissolved O₂ (1.3 x 10⁻⁴ m/kg). Our modeling predicts, as expected, that addition of volcanic gases to these waters results in decreased pH and increased levels of sulfate and chloride in solution; the volcanic gas composition used in these calculations is a restored gas analysis from Kilauea, Hawaii [8], that we modified to contain slightly higher concentrations of HCl than those actually measured.

Chemical Modeling of Rainwater-Basalt Interactions: We modeled the interactions of rainwater and basalts by theoretically titrating increments of an assumed martian basalt composition into the various calculated rainwater compositions. To represent martian basalt, we used the composition inferred by Longhi and Pan [9] for the Chassigny parent magma; lacking reported S, Cl, and P, we assumed for our calculations that this basalt composition contains an additional 165 ppm NaCl and 1375 ppm FeS (the median for terrestrial basalts [4,10]) and 0.8 wt% P₂O₅ (the value inferred for the parent magma of another SNC meteorite [9]).
Modeled reactions of the basalt with acid rainwaters (those acidified by volcanic gases and those under high atmospheric pCO2) initially produce ferric and manganese hydroxides, kaolinite, smectites, chalcedony, and apatite [Ca5(PO4)3(OH, Cl, F)] at high water/rock ratios (>100) and low pH values (<5); the greater the initial acidity, the greater the amounts of basalt converted to these minerals. As progressive rock additions cause the pH to rise to near 8 (at water/rock ratios <10), dolomite and calcite are also predicted to form. Reactions of the basalt with less acidic rainwaters are predicted to form mixtures of ferric hydroxides, apatite, carbonates, and smectites. Ferric hydroxides and ferric smectites are predicted to form under all but the most reducing conditions; such conditions probably existed only very early in martian history [4].

Chemical Modeling of Evaporation: Results of paths modeling the evaporation of rainwaters after reaction with basalted according to the initial sulfate, chloride, and carbonate content of the waters. Those waters acidified by volcanic SO2 prior to reactions with basalt produce gypsum as the dominant sulfate mineral with relatively small degrees of evaporation (less than 0.1 wt% H2O removed); waters that obtain their sulfate solely from dissolution of the basalt precipitate gypsum only after extensive evaporation. Other minerals predicted to form by limited evaporation include dolomite, calcite, potassium feldspar, and sodium feldspar. We are currently modeling the more extreme evaporation conditions needed to produce NaCl, MgSO4, and other soluble salts.

Phosphates in the Martian Regolith: An interesting prediction of our calculations is the formation of apatite during the weathering of basalt containing phosphorus. If aqueous chloride is present in high enough concentrations, the modeling predicts that chlorapatite will form. Our modeling is thus permissive that apatite (or other phosphate phases such as crandallite) could have formed in the martian regolith (1) as an alteration product of basalt and (2) possibly as an evaporation product of phosphate-bearing waters; the extent of phosphate enrichment in the regolith would depend on the primary phosphate content of the basalt, the concentration of phosphate in the waters, and the water/regolith ratio. It is possible that phosphate could partly make up the low totals measured in the Viking analyses [11]. Furthermore, chlorapatite could, if present, serve as a mineralogical host for some of the chloride measured in the regolith.

Summary: Our chemical modeling predicts that weathering of martian basalt in an early, warm, wet climate (with at least some free atmospheric oxygen) would have produced ferric and manganese hydroxides, chalcedony, kaolinite, apatite (or other phosphate minerals), dolomite, and calcite; smectite clays may have also been produced if kinetic barriers to their nucleation were overcome. Acid rain would have enhanced these weathering reactions, and may have provided a source for the high Cl measured in the regolith. Limited evaporation of waters that had previously reacted with basalt leads to formation of gypsum, calcite, dolomite, sodium and potassium feldspars, and some phosphate minerals. Waters initially high in dissolved sulfate and chloride (such as acid rain) require substantially less evaporation to reach saturation with gypsum (and other readily soluble salts) than waters that obtain their sulfate and chloride solely from the rocks. Our modeling also indicates that apatite or other phosphate minerals are likely products of basalt weathering, and should therefore be considered both as possible components of the martian regolith and as possible mineralogical residences for regolith Cl.


INFRARED OPTICAL PROPERTIES OF MARS SOIL ANALOG MATERIALS: PALAGONITES. Ted L. Roush, Department of Geosciences, San Francisco State University, San Francisco CA 94132, USA (also an associate at NASA Ames Research Center, Space Sciences Division, Mail Stop 245-3, Moffett Field CA 94035–1000, USA).

The globally distributed bright soils on Mars represent products of chemical alteration of primary igneous materials. As such, understanding the chemistry and mineralogy of these soils provides clues about the nature of the parent materials and the type, duration, and extent of the chemical weathering environments on Mars. Such clues are key in developing an understanding of the interior and surficial processes that have operated throughout Mars' history to yield the surface as it is currently observed. The generally homogeneous nature of these soils is illustrated by a variety of observational data. These data include (1) direct determination of elemental abundances by the X-ray fluorescence instruments on both Viking landers [e.g., 1–4], (2) Earth-based telescopic observations [e.g., 5–7], and (3) space-based observations [e.g., 8, 9, 10].

Based on their spectral properties in the visible and near-infrared, terrestrial palagonitic soils have been suggested as analogs for the bright regions on Mars [10–12]. Palagonites represent the weathering products of basaltic glass and as such are composed of a variety of minerals/materials. In order to gain an understanding regarding the chemical, mineralogical, and spectral properties of a broad suite of palagonites, several samples were collected from the eastern and central regions of the island of Hawaii. The underlying collection strategy was to select samples from a variety of current climatic environments. As a result, samples were collected from regions that have a wide range in the annual amount of precipitation. Additionally, samples were collected from regions that have similar amounts of annual precipitation. For example, the summit region of Mauna

Fig. 1. Solid line is the initial calculated mass extinction coefficient (c) of one palagonite sample. The dotted line is the straight line continuum and the dashed line is the difference between ε and the continuum.
Kea and the south Point peninsula receive 38 cm and 51 cm of annual rainfall respectively. Because of the proximity of the Pacific Ocean, the south Point region is continuously influenced by spray due to wave action, and additionally is likely to have a higher relative humidity compared to the summit region of Mauna Kea.

Each palagonite sample was dry sieved to the <38-μm grain size fraction. Approximately 0.9–1.3 mg of a palagonite sample was thoroughly mixed with 400 mg of spectroscopic grade KBr in a methanol-powder slurry. After evaporation of the methanol, 200 mg of this mixture was pressed into a pellet using a 12-mm-diameter die at a pressure of 6.89 Kbar for 5 min. Each pellet was placed in the sample beam of a Nicolet 7199 model spectrometer and the transmission was measured at 4 cm⁻¹ spectral resolution over the 400–4000 cm⁻¹ frequency range. This transmission was divided by the transmission of a pure KBr pellet. The thickness of each pellet was measured with a vernier micrometer after data acquisition. To facilitate a comparison of the spectral behavior of these different samples a parameter is required that describes the intrinsic absorptive properties of the materials. One such parameter is the mass extinction coefficient (ε), which is defined as the extinction per centimeter of a gram of material per gram of KBr. From each ratio spectrum ε was calculated via

\[ ε = \frac{-\log_{10}(T)}{ct} \]

where c is the concentration of the sample in g of palagonite per g of KBr, t is the measured thickness of the pellet in cm, and T is the ratio of the transmission of the palagonite pellet to that of the KBr pellet.

Because the scattering properties of the individual pellets can vary substantially, a straight line continuum was fit using the minima points of the ε curves and then subtracted to yield a corrected ε. An example is shown for one sample in Fig. 1. The solid curve is the initially calculated ε, the dashed line is the straight line continuum, and the dotted line is the final ε.

Figure 2 shows ε as a function of frequency for two of the palagonite samples and several extinction maxima are labeled. The broad maxima labeled A and centered around 3400 and 3500 cm⁻¹ is due to the O-H stretching fundamentals of molecular water. The triplet feature labeled B located between 2850 and 3000 cm⁻¹ occurs in the region of the C-H stretching fundamental, although overtone and/or combination modes of a carbonate group may also yield features at these frequencies. Feature C near 1650 cm⁻¹ is due to the H-O-H bending fundamental of molecular water. The feature labeled D, occurring between 1360 and 1490 cm⁻¹, may be due to either C-H stretching modes or the presence of carbonate groups. The two broad remaining features labeled E and F, occurring in the 1300–800 cm⁻¹ and 700–500 cm⁻¹ regions, are both due to Si-O vibrations. The higher frequency feature, E, corresponds to the stretching mode while the lower frequency feature, F, corresponds to the bending mode.

Figure 2 also illustrates the major differences observed in the ε of these palagonite samples. There are two classes of spectra. Class A exhibits a single SiO ε peak at frequencies slightly higher than 1000 cm⁻¹ and an associated ε doublet located between 400 and 600 cm⁻¹. Class B exhibits a single SiO peak at frequencies lower than 1000 cm⁻¹ and an associated singlet near 600 cm⁻¹. It also appears that Class B has a higher ε due to molecular water. Some class A εs also exhibit an indication of structural OH as evidenced by distinctive inflections near 3600 cm⁻¹ and between 870 and 940 cm⁻¹.

The identity of the sources of the major, and less distinctive, differences observed in the ε of these samples awaits independent determinations of the chemistry and mineralogy of these samples. Nevertheless, the results from the current study lead to the conclusion that the infrared properties of palagonitic soils from a wide variety of environments generally fall into two categories. The implication for interpretation of Mars thermal infrared data is that perhaps only a limited number of terrestrial analogs need be considered.

more enhanced by the evaporation of the water. Such saline bodies of water are resistant to freezing, and this ocean could persist as a liquid body on the surface for an extended period of time. There is at present a hypersaline lake in Antarctica (Deep Lake) that never freezes, even though the average temperature there is well below the freezing point of pure water [4].

To study this process, I have developed a geochemical cycle model for the atmosphere-hydrosphere-regolith system of Mars. The treatment of geochemical cycles as complex kinetic chemical reactions has been undertaken for terrestrial systems in recent years with much success [5–8]. This method is capable of elegantly handling the interactions between the simultaneous chemical reactions needed to understand such a system. Geochemical cycles may be represented mathematically by systems of nonlinear differential equations. These systems, which include equations with timescales of widely differing lengths, are considered “stiff.” Such systems require care in the method of integrations, so that a stable solution may be found within a reasonably small time. The present model uses the reverse Euler method, with the computer algorithm based on that presented by Walker [9].

A plot of the decrease in the atmospheric pressure, and subsequent increase in salinity of the ocean, is shown in Fig. 1. Secondary effects of the decrease in water volume, and of the increase in salinity itself, are not included.


**Fig. 1.** Plot of the disappearance of a CO$_2$ atmosphere, and associated increasing alkalinity of the ocean, with time. These results assume no recycling of carbonates back to the atmosphere and no change in salinity.

![Graph](image-url)

**Fig. 2.** Experimental investigation of carbonate formation on Mars. Stuart K. Stephens$^1$, David J. Stevenson$^1$, and Leon F. Keyser$^2$. $^1$Division of Geological and Planetary Sciences, California Institute of Technology, Pasadena CA 91125, USA, $^2$Mail Stop 183-901, Jet Propulsion Laboratory, Pasadena CA 91109, USA.

**Introduction:** If Mars had an early dense CO$_2$ atmosphere (>1 bar), then the transition to the present low pressure (7 mbar) may require carbonate formation without liquid water (e.g., [1], although [2] questions the early greenhouse on the basis that modeled CO$_2$ condensation prevents buildup of such a pressure). Our experiments...
[3] have focused on the kinetics of such “dry” reactions. Booth and Kieffer [4,5] suggested rates consistent with removal of >1 bar of CO$_2$ over geologic time, but their results were questionable since their reactions did not proceed beyond one monolayer of carbonate formed on powdered silicate under martian conditions. Theoretical considerations [1] implied that if nonporous rinds formed, then the rate of carbonate formation would be limited by diffusion, not reaction. Therefore, our experiments optimize reaction conditions to test for continued carbonate production, as well as to verify previous conclusions for initial growth of carbonate.

**Models of Diffusion-limited Carbonate Formation:** Our previous models [3] envisioned carbonate formation at the CO$_2$-silicate interface in the presence of carbonate or bicarbonate ions formed by the interaction of gaseous CO$_2$ with a thin film (monolayer?) of adsorbed OH [5]. Considering a tarnishing reaction where the build-up of product is limited not by reaction rate but by diffusion rate, and using reasonable extrapolations from experimentally measured diffusivities, we derived an equation (dP/dt = f(P)) describing the history of the CO$_2$ atmosphere.

Results of this model (Fig. 1, for two diffusivities) show that the CO$_2$ pressure decays very rapidly in the first few 100 m.y. as a result of higher temperatures and therefore higher diffusivities. More recently than ~2 b.y. ago, the pressure declines gradually—roughly exponentially with a 1-2-b.y. time constant. Buffering effects of CO$_2$ storage in the regolith and polar ice caps [6] are not modeled and would modify our results slightly.

**Present Experiments: Setup.** Our design (Fig. 2) detects reaction of CO$_2$ ( premixed with H$_2$O vapor) with powdered diopside to form carbonate, via careful monitoring of pressure changes in a temperature-regulated environment. Booth and Kieffer [4,5] relied on low-P/T simulations of Mars, whereas we optimize experimental conditions for measurement of a reaction rate that will allow us to extrapolate to martian conditions. (Calculations by Gooding [7] suggest that we are still operating in a thermodynamically favorable regime for the diopside → carbonate reaction.) Also, we use differential pressure measurement in a small volume of atmosphere, instead of relying on acid to dissolve carbonate and infer a reaction [5].

**Fig. 3.**

Initial $P(\text{H}_2\text{O})$: 85 mb (> $P_{\text{sat}}$)
Initial $P(\text{CO}_2)$: 965 mb
T: stages I, III, V, VII, IX -25°C
stage IV -10°C
stages II, VI, VIII, X +20°C
Sample: diopside powder, CaMgSi$_2$O$_6$
mass = 5.0 g
specific surface area = 3 m$^2$/g
of CO₂ loss measured in the sample volume. If this is not a leak or a long-timescale adsorption effect, then it represents a carbonate reaction in the presence of (presumably vapor-phase) water. Since there was a possibility that this -25°C signal represented continued adsorption, we attempted to test this by raising T briefly to -10°C (stage IV), but dPD/dt did not change enough to distinguish adsorption from reaction.

Periods of warmer, wetter conditions (+20°C, liquid water possible) occur in stages II (power out → no data), VI, VIII, and X. The initial rise and fall of PD over 1-2 days is due mainly to the competing effects of (1) PD ~ T in the two volumes and (2) desorption of CO₂. The subsequent rapid rise of PD in stages VIII and X is very interesting. Since long-timescale adsorption of CO₂ should be more vigorous at low temperatures, the most likely explanation for our signal is a chemical reaction that takes CO₂ out of the sample volume.

Discussion. Clearly, something is going on in stage X of this experiment. The role of water is not clear: The amount of H₂O introduced is comparable to the total change in pressure, and that suggests the possibility that water is somehow "used up" (one-on-one with CO₂) in a reaction. However, arguing against that, (1) we do not expect water to be destroyed in a carbonate reaction and (2) the total change in pressure is also comparable to that expected due to formation of a monolayer of carbonate (again, assuming one-on-one with CO₂). We have not yet addressed whether any reaction proceeds beyond one monolayer.

In Fig. 4, we plot dPD/dt vs. t, and find that a model of the form PD = a + bt + c exp(-dt), fits well, especially beyond t ~ 65 days. An exponential form is consistent with a process in which something runs out (e.g., water or available surface area). The fact that we achieve a better fit using values of b > 0 suggests that a roughly linear effect contributes to—but is not responsible for all of—our signal. This effect may represent a leak, or it may represent reaction (possibly involving diffusion through a product layer) beyond the surface reaction that takes place as a monolayer of carbonate is formed. A speculative upper limit can be placed on the diffusivity through this monolayer (D < 10⁻¹² cm²/s).

A rate of b ~ 0.25 mbar/day would correspond to CO₂ storage on Mars rapid enough to get rid of 1 bar in 10⁶ to 10⁷ years (assuming an available specific surface area equivalent to that in a 1-m powdered regolith). This rate is similar to that reported by Booth [5], and is consistent with the diffusion-limited scenario we initially proposed for continued reaction beyond a monolayer [1,3].

Continuing Work: Analysis. If a monolayer of carbonate has formed in the experiment described above, then it represents ~0.3 wt% of our sample, which is barely detectable with reflection IR spectroscopy. We therefore intend to analyze the sample at the end of the current experiment. We also have designed an experiment to subject a fresh surface of diopside (e.g., forsterite-Mg₂SiO₄), greater sample mass, and lower initial P(CO₂) levels. Eventually, we plan to apply experimentally derived reaction rates to models of the past and present CO₂ cycle on Mars (e.g., [8]), which would include the effects of diffusion through a rind, as well as other sources and sinks.


WEATHERING PROCESSES IMPLIED FROM ANALYSIS OF SMALL MARTIAN AVALANCHE CHUTES. R. Sullivan, Department of Geology, Arizona State University, Tempe AZ 85287, USA.

It has been proposed that the smaller features of martian spur and gully slope morphology, located along the upper walls of Valles Marineris, are avalanche chutes [1]. A three-dimensional stability back-analysis technique was developed and applied to these small avalanche chutes, yielding average values of cohesion and angle of internal friction for the mobile layer materials on these slopes at time of each slope failure [2]. Generally, the analysis showed that at the time of each slope failure material strengths had been reduced to those of moderately cohesive debris down through depths of tens of meters. These results have implications and possible constraints for the nature and rate of martian weathering processes.

Dry mass-wasting has long been considered responsible for the recession of the walls of Valles Marineris [1,3,4], but no explanation has been offered as to why steep martian slopes have receded, and may still be receding, forming avalanche chutes, talus, and larger spurss and gullies. This is an important issue, because seemingly older, yet well-preserved features (for example, fluidized ejecta impact craters) are prominent on plateaus adjacent to receding/receded scarps. If older topographic features exist on adjacent plateaus in a relatively fresh state, then the clear evidence of slope recession in the intervening time, including prominent talus production, seems paradoxical. The key question is, after all, why should martian steep slopes recede at all? Recession of such prominent slopes represents profound physical change to the martian landscape. Why should a combination of degradational processes affect one set of features—the steep slopes—and not older features nearby?

One possibility is that weathering (and associated weakening) of martian surface materials occurs broadly across the martian surface—not just on steep, receding slopes—but this process or suite of processes is revealed especially on steep, receding slopes by the periodic removal of weathered layers under the influence of gravity. Debris or soils generally have significantly lower strengths than the
sublimation, carries the binding agent (ice) away into the atmosphere. The sublimation of subsurface ice is an appealing hypothesis in some respects, but the recent presence of subsurface ice in equatorial latitudes is an unsettled question [5] and direct evidence for the existence or nonexistence of subsurface ice (at depths appropriate to this issue) will probably be a long time in coming.

As an alternative to invoking the unproven presence of a volatile ice in the martian near-subsurface, how could our present knowledge of the current surface and near-subsurface environment and its suite of weathering processes explain the geomorphological observations and stability analysis results? This is a difficult task because our knowledge of possible chemical weathering reactions, reaction efficiencies, and products is relatively unconstrained. As yet unmeasured or unimagined characteristics of the martian subsurface environment and its weathering reactions may be responsible. The results of stability back-analysis may offer some useful constraints in addressing this question in the future, however. Figures 1 and 2 represent avalanche chute analysis results plotted such that the average indicated cohesion and angle of internal friction for each avalanche chute mobile layer is plotted vs. the depth (normal to the surface) of each chute floor. A crude "soil profile" of cohesion and angle of internal friction results, and is presented here as a possible constraint for the future analysis of martian weathering processes and products. It should be emphasized that Figs. 1 and 2 are only suggestive and should be used with caution, for together they do not represent a true soil profile from a single location, but are composed of results from 25 different locations and slope angles, and only where slope failure has occurred. For instance, it is not implied here that the stronger materials at the deeper points weather upward to the weaker materials at the points nearer the surface; initial strength variations of pristine parent material from one location to another could account for much of the variation in strength of these presumably weathered products. Nevertheless, these results should be suggestive of the range of cohesion and angle of internal friction values of martian subsurface materials to be found in the study areas. Figure 1 shows that where these avalanche chutes have formed, cohesion increases gradually with depth. Figure 2 shows that angle of internal friction does not vary strongly with depth for the analyzed chutes (as might be expected, for angle of internal friction is not diagnostic for most classes of geological materials). The slight increasing trend with depth in Fig. 2 is probably an artifact; the shallowest chutes were only visible because their average slope angles were shallow, making their headscars more prominent, and shallow slope angles lead to smaller indicated angles of internal friction in the calculations.


![Fig. 2. Angle of internal friction vs. depth for 25 martian avalanche chutes.](image-url)
The Lafayette achondrite, one of the nakhlites of probable martian origin, is an igneous rock consisting mostly of augite and olivine, with interstitial feldspar, sulfides (pyrrhotite), high-Si glass, and other phases [1–3]. Like Nakhla itself, Lafayette contains veinlets of hydrous alteration materials [1–7].

Sample and Methods: We studied thin sections of sample ME2116 (Field Museum, Chicago), using scanning and transmission electron microscopy (SEM and TEM) with energy dispersive X-ray spectrometry (EDS) as in [4,8]. SEM elemental analyses were done in spot mode (1 μm) and in rastered selected area mode (to 10 μm); TEM elemental analyses were for areas from 100 to <10 nm diameter. Standards were well-characterized minerals; Na abundances in SEM/EDS were corrected for interference from FeL X-rays [7]. SEM/EDS analyses of olivine in Lafayette are nearly identical to EMP analyses from [1–3].

Petrography: Rusty veinlets of aqueous alteration material replace olivine, pyroxene, and high-Si glass, and occur as intergranular films. Olivine is replaced by veinlets and spikes of rhombic cross sections [7]. The veinlets commonly have exterior zones of 3–5-μm phyllosilicate grains, cut by veinlets of <100-nm material ± 100-nm grains of high secondary electron (SEI) brightness. These fine-grained veinlets may cross-cut each other, and commonly have cores with higher SEI brightness than the rims. By TEM, the veinlets consist of felted masses of smectite (platy or fibrous grains, 1.0-nm lattice fringes, composition consistent with trioctahedral clay) and iron oxides, probably hematite and ferrihydrite. These materials are cut by later veinlets consisting of magnetite and void space.

Preterrestrial Origin: These aqueous alteration materials are preterrestrial, as shown by changes in the alteration veinlets as they approach the fusion crust [6,7]. More than 0.5 mm from the fusion crust, veinlets are as above. Between 0.5 mm and 0.3 mm, the veinlets are as above, and become depleted in volatile elements. Within 0.3 mm of the crust, the veinlets are melted to uniform vesicular glass and lose almost all volatiles. The melted veinlets are cross-cut by the fusion crust. These data suggest that the veinlets existed before Lafayette entered the Earth’s atmosphere, and that the veinlets were heated, devolatilized and melted by passage through the Earth’s atmosphere [3]. These inferences are consistent with the oxygen isotope data of Karlsson et al. [9].

Composition: The compositions of the alteration materials are consistent with their mineralogy as inferred from SEM and TEM [6,7]. The rough correlation of K and Al is consistent with smectite. Abundances of Cl and S are correlated for analyses with less than 1% SO₃ and 0.5% Cl; the remaining analyses are richer in SO₃ or Cl. The latter analyses suggest discrete S-rich (sulfate) and chloride phases similar to those in Nakhla [4,5]. Correlation of S and Cl at lower abundance levels suggests a single carrier phase, perhaps ferricyanide acting as an ionic sorbent.

Among the analyses, abundances of Fe and Si are inversely correlated, and are collinear with the compositions of the olivine and high-Si glass (Fig. 1a). A similar collinearity is seen for Mn and Fe. Imperfect collinearities are seen for abundances of Mg, Al, and Fe (Fig. 1b).

Geochemistry: The alteration veinlets here and in the other SNC meteorites are the only available traces of the hydrosphere on their parent planet, presumably Mars. As such, they may have significant implications for the origins, cycling, and availability of water on Mars. Detailed studies are in progress, and we can here point out a few features of the geochemistry of Lafayette’s alteration.

Temperature: Lafayette was altered at low temperatures, almost certainly below 100°C. This limit comes from the presence of ferrihydrite in the alteration materials. Ferrihydrite dehydrates read-

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Fig. 1. Chemical relationships among Lafayette’s alteration (SEM/EDS analyses). (a) FeO-SiO₂. Note that alteration materials are collinear with olivine and high-Si glass. (b) MgO- and Al₂O₃-FeO. Average compositions of materials in Lafayette: MgO, squares; Al₂O₃, triangles. Note that alteration materials in a phase are enriched in the essential constituents of that phase (Mg in olivine; Al in glass). Note also that the average composition of all alteration materials is closely approximated by a mixture, the lines, of olivine and high-Si glass (+ water).
ily to hematite or goethite at temperatures above 100°C [10,11]. On Earth, ferricyanide and ferrous smectite are most commonly produced during the weathering or low-temperature alteration of basic rocks [12-14], notably weathering in the dry valleys of Antarctica [15,16]. The porous oxide alteration materials must also have formed at low temperature; otherwise, ferricyanide in the other alteration materials would have been converted to other minerals. At such temperatures, the alteration phase was almost certainly liquid water.

**Oxidation state.** The oxidation state of Lafayette's alteration was near or below that of the magnetite-hematite oxygen buffer, distinctly more reducing than the alteration of Nakhl [4,5] or the martian atmosphere. This estimate of oxygen fugacity comes from the presence of magnetite (or maghemite) and ferricyanide in the alteration materials. Because the assemblage magnetite-hematite-ferricyanide-water is invariant in Fe-O-H (constant pT), the oxygen fugacity buffered by magnetite-ferricyanide-water must be below that of magnetite-hematite, but probably not by much.

**Fluid composition and transport.** The compositions of the altering fluids are poorly constrained so far. The fluids did contain some dissolved salts, as shown by the moderate abundances of S, Cl, and P in the alteration materials. The S and Cl may be present in part as minute grains of discrete phases (see above), not only as adsorbed anions. Note that the Lafayette's magmatic sulfides and phosphates could be the sources of the S and P in the alteration materials.

In a similar vein, it appears that the altering solutions transported little into or out of Lafayette. Figure 1 shows the collinearity of FeO vs. SiO₂ for the alteration materials, olivine, and high-Si glass. The collinearity suggests that the Fe-Si-"others except water" abundances of the alteration material can be explained merely by mixing of Lafayette olivine and high-Si glass (± iron oxides). Limited chemical transport within Lafayette is suggested by MgO and Al₂O₃ concentrations in the alteration materials from different settings. In Fig. 1b, note that alteration materials in olivine are relatively rich in MgO, just as is the olivine, and that alteration materials in mesostasis glass are relatively rich in Al₂O₃, just as is the glass. This chemical similarity between host and alteration product suggests that Mg and Al were not completely mobile during Lafayette's alteration. For both elements, the average composition of all alteration materials lies close to the line between olivine and the mesostasis glass.

So there is no evidence that major elements in the alteration materials have been transported into or out of Lafayette during alteration (Fig. 1). In effect, alteration of Lafayette was a water-mediated reaction between its own olivine, Si-rich glass, and possibly other (S-, Cl-bearing) phases. The altering solutions need have transported little except water, Cl, and possibly S. Lacking evidence for extensive chemical transport, it seems likely that water-rock ratios during alteration were low.

**Geological setting.** From the limited data available, Lafayette probably was altered in a geological setting of episodic water flow or injection. Long-term saturation with aqueous fluid, as beneath a water table, does not seem consistent with available data as we understand them.

First, it seems unlikely that Lafayette was saturated with aqueous solutions for long times, because such soaking ought to produce alteration materials on all surfaces and cracks that expose olivine or high-Si glass [17]. In fact, the alteration materials are present only along some cracks and some grain boundaries, and not pervading the rock.

Direct evidence for episodic alteration comes from the presence of different varieties of alteration materials in cross-cutting relationships, and from the cross-cutting relationships between different veinlets of fine-grained alteration materials. Episodic alteration is also consistent with the presence of alteration materials only adjacent to cracks, which (as above) suggest fluid flow through the rock and not stagnant fluid in the rock.

**Setting on Mars.** We cannot specify a geological environment for the aqueous alteration of Lafayette. Possible martian causes for episodic, limited alteration of basaltic rocks may include melting of permafrost by magma injection [18], hydrothermal convection cells (distal only) [19], and thermal transients associated with meteoric impacts [20]. Given the limited data on the martian hydrosphere, this list is likely not exhaustive.

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**ATTEMPTS TO COMPREHEND MARTIAN SURFACE PROCESSES THROUGH INTERPRETATION OF THE OXYGEN ISOTOPIC COMPOSITIONS OF CARBONATES IN SNC METEORITES.** I. P. Wright¹, C. T. Pillinger¹, and Monica M. Grady², ¹Department of Earth Sciences, Open University, Walton Hall, Milton Keynes MK7 6AA, UK, ²Department of Mineralogy, The Natural History Museum, Cromwell Road, London SW7 5BD, UK.

The SNC meteorites are known to contain trace quantities of a variety of secondary minerals such as carbonates [e.g., 1,2], sulfates [e.g., 3,4], and aluminosilicates [e.g., 4]. Since these constituents are thought to be mostly preterrestrial in origin, their study has the potential to provide rigorous constraints on the nature of martian weathering processes. However, this line of investigation is potentially complicated by the presence within the meteorite samples of any additional weathering products produced by terrestrial processes. Examination of such terrestrial components is important since weathering processes that affect meteorite samples following their fall to Earth might have some bearing on the nature of analogous processes at the surface of Mars. It is obviously necessary to try and distinguish which secondary components in SNC meteorites are terrestrial in origin from those that are preterrestrial. Herein consideration is made of the stable isotopic compositions of weathering products in two SNC meteorites: EET A79001 (a sample collected from Antarctica) and Nakhl (a fall from Egypt, 1911).

Carbonate minerals in EET A79001 and Nakhl have similar carbon and oxygen stable isotopic compositions [1,5,6], suggestive of a common (martian) origin. Nonetheless, it has recently been
reported that carbonates in EET A79001 contain high levels of $^{14}$C [7], 57-95% of which might be expected if the minerals were of modern terrestrial origin. In contrast, the carbonates in Nakhla contain very little $^{14}$C [8], which endorses a preterrestrial origin for these minerals. It has previously been pointed out that the stable isotopic data for the carbonates in EET A79001 ($^{13}$C = 7+10‰, $^{18}$O = 21‰) and Nakhla ($^{13}$C = 6+9‰, $^{18}$O = 23 to 26‰) are somewhat unusual compared to common terrestrial occurrences of these minerals [5], although it is now clear that there is a similarity with Antarctic weathering products [7,9,10]. While an origin in the Antarctic is clearly possible for the carbonates in EET A79001 this is not the case for Nakhla. It could be argued that in light of the possibilities that EET A79001 has been altered on Earth, the study of this meteorite is ill-advised. However, EET A79001 is an important sample for many reasons—for instance, this sample contains trapped martian atmospheric gases [e.g., 11,12] including CO$_2$ [1,13]. Furthermore, EET A79001 is a fairly large find and is therefore readily available for study. Rather than simply discounting the EET A79001 sample, several in-depth investigations have been carried out in order to try and understand the nature of the carbon compounds in this meteorite [e.g., 5,14].

It remains to be explained why EET A79001 and Nakhla contain carbonate minerals with similar stable isotopic compositions, but which apparently formed on different places (Earth and Mars respectively). There are a number of possible explanations—for example, EET A79001 could contain (1) martian carbonates similar to those in Nakhla, but with the high levels of $^{14}$C incorporated as a result of poorly understood exchange process(es) acting on Earth; (2) secondary minerals produced in the Antarctic, with $^{13}$C and $^{18}$O not dissimilar to the carbonates in Nakhla, in addition to martian carbonates; (3) only preterrestrially produced carbonates, the similarity in stable isotopic compositions to Nakhla being due to the operation of similar processes at the surface of Earth (i.e., in Antarctica) and Mars.

Point (1) has been addressed briefly by Jull et al. [7]—the carbonates in EET A79001 may originally have been preterrestrial, but their isotopic compositions could have subsequently been reset by fluid-related processes in Antarctica. This hypothesis can only be assessed by further studies. The notion that EET A79001 contains two different sorts of carbonate, option (2), is demonstrably true [2] and attempts have been made to determine the carbon isotopic compositions of the different varieties [14,15]. In regard of point (3), the martian surface environment can undoubtedly be considered similar in some respects to Antarctica. As such it is instructive to know more about how Antarctic weathering products may be formed.

When meteorites are subjected to weathering processes in Antarctica, a fluid phase, which derives from the ice, is involved. For illustrative purposes it can be assumed that Antarctic ice has a $^{18}$O value of about -40‰ (compared to 0‰ for terrestrial ocean water)—the depletion in $^{18}$O is a reflection of the temperature of precipitation of snow, etc., that compacts to form the ice sheets. An assessment of the oxygen isotopic compositions of Antarctic weathering products is obtained from measurements of carbonates associated with a variety of ordinary chondrites collected from Antarctica. The carbonates are found to have $^{18}$O in the range -5 to +25‰ [16,17]. One particular Antarctic meteorite (LEW 85320) has been studied in detail due to the extensive deposits of weathering products that cover the sample’s exterior surface—these have $^{18}$O values between +9 and +22‰ [9,10]. Although Antarctic weathering products can take on many different forms, since the carbonates in SNC meteorites are ascribed to calcite [2,18], it is convenient to consider only this mineral in the following discussion. As noted above, carbonates in EET A79001 have $^{18}$O of +21‰, while those in Nakhla have $^{18}$O of +23 to +26‰. At 0°C, $^{18}$O of calcite produced in isotopic equilibrium with Antarctic melt water would be about -5‰; in order to obtain a $^{18}$O of +25‰, carbonate would have to be precipitated at an unrealistic temperature of -70°C. Thus, while it may be possible, in theory, to derive some of the Antarctic weathering products observed in meteorites by exchange with, or formation from, Antarctic melt water, it is not possible to explain the oxygen isotopic data for carbonates in EET A79001 entirely by this mechanism. It may therefore be argued that the oxygen isotopic composition of the melt water has become enriched in $^{18}$O by exchange with either the silicates of the meteorite ($^{18}$O of +44‰) or atmospheric CO$_2$ ($^{18}$O = +41‰). Isotopic exchange with the meteorite itself is unlikely in this instance since this would depress the $^{18}$O of the silicates [e.g., 19], a feature not observed in the case of EET A79001, which has $^{18}$O similar to all other SNC meteorites [20,21]. Calcite grown in isotopic equilibrium with atmospheric CO$_2$ would have $^{18}$O of +29‰. It is apparent that the range of $^{18}$O values displayed by Antarctic weathering products (-5 to +25‰) demands the involvement of melt water and atmospheric CO$_2$. If the carbonates in EET A79001 are also Antarctic weathering products then these too must have been formed by a similar process.

It is now possible to equate what is observed on Antarctica to potential processes on Mars. In simplistic terms, if the temperature of carbonate formation is similar to that in operation in Antarctica, then the isotopic constraints can theoretically be satisfied by a martian water phase that has $^{18}$O of -40‰ and atmospheric CO$_2$ with $^{18}$O of +41‰. However, the situation is a little more complicated and notice needs to be taken of the magnitude of the CO$_2$/H$_2$O ratio [22]. The effects of martian weathering can be explored further if the $^{18}$O values of the surface volatiles on Mars are known. This subject has been dealt with extensively by Jakosky [23]—in brief, the $^{18}$O of atmospheric CO$_2$ has been measured at 0 ± 5‰, while $^{18}$O of atmospheric water vapor is -100 ± 60‰. Unfortunately these measurements are not particularly useful for constraining the operation of martian surface weathering processes because of their large associated uncertainties. A recent study of water liberated from SNC meteorites by heating [24] has shown coupled variations in $^{18}$O and $^{18}$O that indicate that at least some of the SNC meteorites (Nakhla included) contain water of preterrestrial origin. The exact form of this water is unknown, but SNC meteorites are known to contain hydrated minerals of both high temperature [e.g., 25] and secondary origin [e.g., 18]. Since variations in $^{18}$O of $^{18}$O indicate that the hydrated phases of SNC meteorites are not in isotopic equilibrium with the silicates, it seems logical to conclude that the majoritic of the water in SNC meteorites arises from secondary weathering products. Water released from Nakhla was about 10‰ enriched in $^{18}$O compared to a “terrestrial control sample”; on the other hand, water from EET A79001 had a similar $^{18}$O to the control [24]. Unfortunately the $^{18}$O of the control is not given in [24], but if it is assumed that it was a rock of Antarctic affinities then it is possible to accept that the water in EET A79001 was imparted by terrestrial weathering processes. In other words, the water in EET A79001 arises from minerals produced in equilibrium with Antarctic melt water ($^{18}$O = -40‰) and as such would have $^{18}$O values of about -10‰ (there being an ~30‰ oxygen isotopic fractionation at 0°C for a number of common hydrated minerals). Taking this further, it is apparent that $^{18}$O of hydrated minerals in Nakhla would be about 0‰; if these were formed in isotopic equilibrium with water at 0°C, the $^{18}$O of this water would be ~30‰, which is outside the range of possible values for mean atmospheric water vapor (i.e., $^{18}$O = -100 ± 60‰). If, as seems more likely, the $^{18}$O of the control used in [24] was
somewhere in the range $+10 \pm 5\%$, the hydrous phases of Nakhla would have $\delta^{18}O$ of $-20\%$ and the water from which they were produced would have $\delta^{18}O$ of $-10\%$.

Intuitively, since there is an effective cycling of water vapor through the atmosphere and regolith on Mars, it would be anticipated that any surface fluids (which could induce weathering) would be isotopically similar to the atmospheric water vapor. As such, the disparity noted here is somewhat surprising. However, it has been proposed that the $\delta^{18}O/\delta^{16}O$ values of water released from SNC meteorites were established as a result of isotopic disequilibrium between the lithosphere and hydrosphere of Mars [24]. Perhaps we should conclude that this disequilibrium extends to the near-surface fluids and atmospheric gases on Mars. On the basis of the (admittedly meagre and somewhat incomplete) SNC data it would seem unlikely that surface water on Mars could have $\delta^{18}O$ of $-40\%$. In contrast, if the water has $\delta^{18}O$ of $-10\%$ then carbonates that precipitate in equilibrium with this will have, at $0^\circ C$, $\delta^{18}O$ of $+25\%$, a value commensurate with what is observed in SNC meteorites.


**TABLE 1.** Explanations for Viking results from the literature.

<table>
<thead>
<tr>
<th><strong>GEX release of O$_2$ upon humidification.</strong></th>
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<tbody>
<tr>
<td>KO$_2$</td>
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<tr>
<td>ZnO$_2$</td>
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<tr>
<td>CaO$_2$ in soil</td>
</tr>
<tr>
<td>MnO$_2$</td>
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<tr>
<td>O$_2$ trapped in micropores</td>
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<tr>
<td>Frost on olivine/pyroxene</td>
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<tr>
<td>Amboy Soil under plasma</td>
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<tr>
<td>Oxidized halides</td>
</tr>
</tbody>
</table>

**• LR decomposition of added nutrient.**

| H$_2$O$_2$ | [10] |
| --- |
| Peroxidant (NOO) | [4] |
| Smectite clays | [8] |

**• Lack of organic in martian soil.**

| UV + TiO$_2$ | [13] |
| Dust electrostatics | [14] |
| Ferroxoxy (3-Fe-OH) | [16] |

*See references.
Although H₂O does not appear on the lefthand side of these equations because there is a net production of H₂O in the reaction, it is a necessary catalyst to rapid decomposition reactions. In addition, acids of Br and I can be identified that may play a role in H₂O decomposition, although their stability under GEx sterilization conditions has not yet been determined.

A possible compound for the LR oxidant is H₂O₂, catalyzed by the presence of iron in the soil. The concentration of H₂O₂ required to explain the LR results is about 1 ppm by mass.

**Measurement on Future Missions:** An *in situ* investigation of the putative regolith oxidant is indicated for precursor missions leading to human exploration.

The traditional variable for characterizing the oxidation state of terrestrial soils is Eh. Eh is inappropriate to characterize of the martian regolith for a variety of reasons. For example, Eh is used to determine aqueous speciation and saturation levels for terrestrial soil solutions. In cases where a soil solution exists, successful measurements of Eh can be made if it can be established that the solution is truly in equilibrium with the soil [20]. These techniques are applied with great success in saturated soils, such as rice paddies or long-lived bodies of soil solution. However, in cases where there is no soil solution, the technique is highly suspect.

There are two requirements for successful characterization of the oxidation potential of a soil solution via this method that apply to terrestrial soils, and a third requirement that applies rather uniquely to Mars. First, there must be dissolution equilibrium between the soil and the solution. This ensures that the potential developed at the electrode-solution interface reflects the properties of the soil. In dry terrestrial soils, it is often impossible to determine whether dissolution equilibrium exists, except by maintaining the sample in contact with the solvent/electrolyte for extended periods of time. Even more stringent is the requirement that electrochemical equilibrium between all redox couples exposed to the solution exist. This includes species that may not be soluble in the solution but may exist on the surfaces of the soil grains. Electrochemical equilibration is notoriously slow in terrestrial soils, in spite of the fact that most electrochemical reactions are catalyzed by microorganisms in the soil. In sterile soils, equilibration may be expected to take months; this requirement severely complicates implementation of an Eh measurement on a landed experiment. Even more problematic is the requirement that, if no soil solution is present, a solvent must be added in order to make the Eh measurements. In terrestrial soils, H₂O is the solvent of choice because of its ubiquity. However, it is implicit that the solvent/electrolyte must be electrochemically neutral with respect to the soil sample. There is abundant evidence, as described above, that the putative oxidant has an electrode potential greater than that of the H₂O₂-O₂ couple, and reacts vigorously with H₂O. Therefore, H₂O is an inappropriate solvent/electrolyte. Addition of H₂O to the soil is likely to destroy the species that we are most interested in measuring. It is possible to substitute another electrolyte with an oxidation potential greater than that of the H₂O₂-O₂ couple. Although the putative oxidant may be electrochemically inert with respect to such a solvent, acetonitrile for example, the dissolution equilibrium requirement then becomes problematic.

A possible approach to measurement of the putative LR oxidant is a specific search for H₂O₂ in the martian soil. We have proposed a H₂O₂ fluorometer, in which an organic acid dimerizes in the presence of H₂O₂. The dimer can be excited by UV radiation, and fluoresces at 400 nm. The fluorescence is directly proportional to the catalytic activity of the H₂O₂ on the acid monomer.

**References:**
